

CLAIMS

We claim:

1. A biochip comprising a solid substrate comprising an array comprising:
 - a) at least one capture probe substantially homologous to a portion of the sense strand of a nucleic acid encoding CPY1A1;
 - b) at least one capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CPY1A2;
 - c) at least one capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CPY1B1;
 - d) at least one capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CPY2C19;
 - e) at least one capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CPY2D6;
 - f) at least one capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CPY2E1; and
 - g) at least one capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CPY3A4.
2. A biochip according to claim 1 wherein said first portion of said sense strand of said nucleic acid encoding CPY2D6 is adjacent to a single nucleotide polymorphism (SNP) position of interest.
3. A biochip according to claim 1 wherein said first portion of said sense strand of said nucleic acid encoding CPY2D6 includes at a terminus a single nucleotide polymorphism (SNP) position of interest.
4. A biochip according to claim 1 wherein said array further comprises at least one capture probe substantially homologous to a portion of the antisense strand of a nucleic acid encoding a protein selected from the group consisting of CYP1A1; CYP1A2; CYP1B1; CYP2C19; CYP2D6; CYP2E1 and CYP3A4.

5. A biochip according to claim 1 wherein said array further comprises at least one capture probe substantially homologous to a portion of a CYP pseudogene.
6. A biochip according to claim 1 wherein said solid support is selected from the group consisting of glass, plastic, ceramic, and PC board.
7. A biochip according to claim 1 wherein said array comprises an array of electrodes.
8. A biochip according to claim 1 wherein said array comprises an array of polymer gel pads.
9. A method of determining the identification of a nucleotide at a detection position in at least one target sequence selected from the group consisting of CYP1A1, CYP1A2, CYP1B1, CYP2C19, CYP2D6, CYP2E1 and CYP3A4, said method comprising:
 - a) providing an array comprising:
 - i) at least one first capture probe substantially homologous to a first portion of a nucleic acid encoding CYP1A1, wherein said first capture probe is directly adjacent to or includes at its terminus a detection position;
 - ii) at least one second capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CYP1A2, wherein said second capture probe is directly adjacent to or includes at its terminus a detection position;
 - iii) at least one third capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CYP1B1, wherein said third capture probe is directly adjacent to or includes at its terminus a detection position;
 - iv) at least one fourth capture probe substantially homologous to a first portion of the sense strand of a nucleic acid encoding CYP2C19, wherein said fourth capture probe is directly adjacent to or includes at its terminus a detection position;
 - v) at least one fifth capture probe substantially homologous to a first

portion of the sense strand of a nucleic acid encoding CPY2D6,
wherein said fifth capture probe is directly adjacent to or includes at
its terminus a detection position;

vi) at least one sixth capture probe substantially homologous to a first
portion of the sense strand of a nucleic acid encoding CPY2E1,
wherein said sixth capture probe is directly adjacent to or includes at
its terminus a detection position; and

vii) at least one seventh capture probe substantially homologous to a
first portion of the sense strand of a nucleic acid encoding CPY3A4,
wherein said seventh capture probe is directly adjacent to or includes
at its terminus a detection position;

b) hybridizing at least one target sequence to its corresponding capture probe to form a
hybridization complex;

c) adding a polymerase and at least one dNTP comprising a label, under conditions
whereby if said dNTP is perfectly complementary to a detection position, said dNTP is
added to a capture probe to form an extended probe;

d) determining the nucleotide at the interrogation position of said extended probe.

10. A method of determining the identification of a nucleotide at a detection position in a target
sequence comprising:

a) providing an array comprising:

i) a solid support with a first surface comprising a hydrogel layer
comprising an array of capture probes;

b) hybridizing said target sequence to at least one of said capture probes to form a
hybridization complex; and

c) determining the nucleotide at said detection position.

11. A method of determining the identification of a nucleotide at a detection position in a target
sequence comprising:

a) providing a solid support with a first surface comprising at least one non self-extension
probe wherein self-extension of said non self-extension probe does not occur in the absence

of said target and wherein, said non self-extension probe includes an interrogation nucleotide;

b) hybridizing said target sequence to said non self- extension probe to form a hybridization complex;

c) contacting said surface with:

i) an extension enzyme; and

ii) at least one chain terminating nucleotide comprising a hapten;

under conditions whereby if said chain terminating nucleotide is perfectly complementary to the base of the target sequence immediately adjacent to the 3' end of said non self- extension probe in the hybridization complex, said chain terminating nucleotide is added to said non self-extension probe to form a modified extension probe;

d) contacting said modified extension probe with the binding partner of said hapten, wherein said hapten is labeled; and

e) detecting the presence of said label to determine the nucleotide at said detection position.

12. A method according to claim 11 wherein said interrogation nucleotide in said non self-extension probe is within two bases of its 3' terminal end and wherein, said 3' terminal end nucleotide is non-complementary to a corresponding base when a self-hybridizing structure of said non self-extension probe is formed.

13. A method according to claim 12 wherein said interrogation nucleotide is the 3' terminal nucleotide.

14. A method according to claim 12 wherein said interrogation nucleotide is the penultimate 3' terminal nucleotide.

15. A method according to claim 11 wherein said non self- extension probe comprises at least two modified nucleotides.

16. A method according to claim 15 wherein said modified nucleotides are exo-cyclic amine modified bases.
17. A method according to claim 15 wherein said modified nucleotides are terminator bases.
18. A method according to claim 16 wherein said exo-cyclic amine modified bases are selected from a group consisting of 2-thio thymine, 2-amino adenine, amine modified cytosine and amine modified guanine.
19. A method according to claim 17 wherein said terminator base is 4-methylindole .
20. A method according to claim 15 wherein said modified nucleotides alter protein binding and are present in the stem region of said non self-extension probe.
21. A method according to claim 20 wherein said modified nucleotide comprises a sugar modification.
22. A method according to claim 20 wherein said modified nucleotide comprises a phosphate modification.
23. A method according to claim 22 wherein said phosphate modifications are selected from a group consisting of phosphorothioates, phosphoramidates, methyl phosphonates, methyl phosphates, H-phosphonates.
24. A method according to claim 11 wherein self-extension of said non self-extension probe is inhibited by short complementary oligonucleotides.
25. A method of determining the identification of a nucleotide at a detection position in a target sequence comprising:

- a) amplifying the target DNA using random primers to generate DNA amplicons;
 - b) transcribing said DNA amplicons to generate RNA target sequences (*in vitro* transcription);
 - c) providing a solid support with a first surface comprising at least one extension probe wherein said extension probe includes an interrogation nucleotide;
 - b) hybridizing said RNA target sequence to said extension probe to form a hybridization complex;
 - c) contacting said surface with:
 - i) a modified reverse transcriptase; and
 - ii) at least one chain terminating nucleotide comprising a hapten;
- under conditions whereby if said chain terminating nucleotide is perfectly complementary to the base of the target sequence immediately adjacent to the 3' end of said non self- extension probe in the hybridization complex, said chain terminating nucleotide is added to said non self- extension probe to form a modified extension probe;
- d) contacting said modified extension probe with the binding partner of said hapten, wherein said binding partner is labeled; and
 - e) detecting the presence of said label to determine the nucleotide at said detection position.

- 26. A method according to claim 25 wherein said modified reverse transcriptase only extends extension probes bound to RNA.
- 27. A method according to claim 11 or 25 wherein said hapten is biotin.
- 28. A method according to claim 11 or 25 wherein said binding partner is streptavidin.
- 29. A method according to claim 11 or 25 wherein said binding partner is Alexa dye labeled streptavidin.

30. A method of determining the identification of a nucleotide at a detection position in a target sequence comprising:

a) providing a solid support with a first surface comprising a solid support with a first surface comprising a hydrogel layer comprising at least one non self-extension probe, wherein self-extension said non self-extension probe does not occur in the absence of said target and wherein, said non self-extension probe includes an interrogation nucleotide;

b) hybridizing said target sequence to said non self-extension probe to form a hybridization complex;

c) contacting said surface with:

i) an extension enzyme; and

ii) at least one chain terminating nucleotide comprising a hapten;

under conditions whereby if said chain terminating nucleotide is perfectly complementary to the base of the target sequence immediately adjacent to the 3' end of said non self-extension probe in the hybridization complex, said chain terminating nucleotide is added to said non self-extension probe to form a modified extension probe;

d) contacting said modified extension probe with the binding partner of said hapten, wherein said binding partner is labeled; and

e) detecting the presence of said label to determine the nucleotide at said detection position.

31. A method according to claim 30 wherein said interrogation nucleotide in said non self-extension probe is within two bases of its 3' terminal end and wherein, said 3' terminal end nucleotide is non-complementary to a corresponding base when a self-hybridizing structure of said non self-extension probe is formed.

32. A method according to claim 31 wherein said interrogation nucleotide is the 3' terminal nucleotide.

33. A method according to claim 31 wherein said interrogation nucleotide is the penultimate 3' terminal nucleotide.

34. A method according to claim 30 wherein said non self- extension probe comprises at least two modified nucleotides.
35. A method according to claim 34 wherein said modified nucleotides are exo-cyclic amine modified bases.
36. A method according to claim 34 wherein said modified nucleotides are terminator bases.
37. A method according to claim 35 wherein said exo-cyclic amine modified bases are selected from a group consisting of 2-thio thymine, 2-amino adenine, amine modified cytosine and amine modified guanine.
38. A method according to claim 36 wherein said terminator base is 4-methylindole .
39. A method according to claim 34 wherein said modified nucleotides alter protein binding and are present in the stem region of said non self-extension probe.
40. A method according to claim 39 wherein said modified nucleotide comprises a sugar modification.
41. A method according to claim 39 wherein said modified nucleotide comprises a phosphate modification.
42. A method according to claim 41 wherein said phosphate modifications are selected from a group consisting of phosphorothioates, phosphoramidates, methyl phosphonates, methyl phosphates, H-phosphonates.
43. A method according to claim 30 wherein self-extension of said non self-extension probe is inhibited by short complementary oligonucleotides.

44. A method of determining the identification of a nucleotide at a detection position in a target sequence comprising:
- a) amplifying the target DNA using random primers to generate DNA amplicons;
 - b) transcribing said DNA amplicons to generate RNA target sequences (*in vitro* transcription);
 - c) providing a solid support with a first surface comprising at least one extension probe wherein said extension probe includes an interrogation nucleotide;
 - b) hybridizing said RNA target sequence to said extension probe to form a hybridization complex;
 - c) contacting said surface with:
 - i) a modified reverse transcriptase; and
 - ii) at least one chain terminating nucleotide comprising a hapten;
- under conditions whereby if said chain terminating nucleotide is perfectly complementary to the base of the target sequence immediately adjacent to the 3' end of said non self- extension probe in the hybridization complex, said chain terminating nucleotide is added to said non self- extension probe to form a modified extension probe;
- d) contacting said modified extension probe with the binding partner of said hapten, wherein said binding partner is labeled; and
 - e) detecting the presence of said label to determine the nucleotide at said detection position.
45. A method according to claim 44 wherein said modified reverse transcriptase only extends extension probes bound to RNA.
46. A method according to claim 30 or 44 wherein said hapten is biotin.
47. A method according to claim 30 or 44 wherein said binding partner is streptavidin.
48. A method according to claim 30 or 44 wherein said binding partner is Alexa dye labeled

streptavidin.

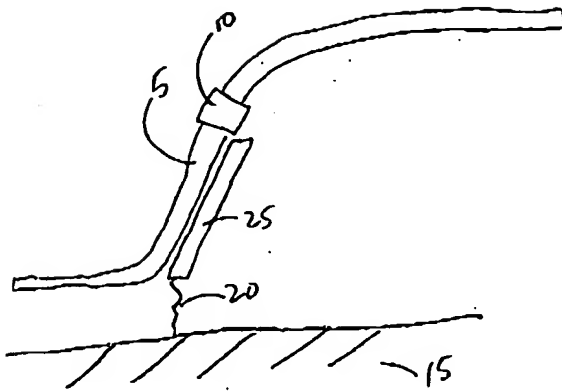


Fig 1A

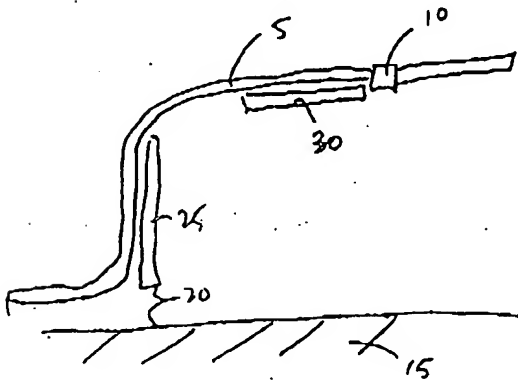
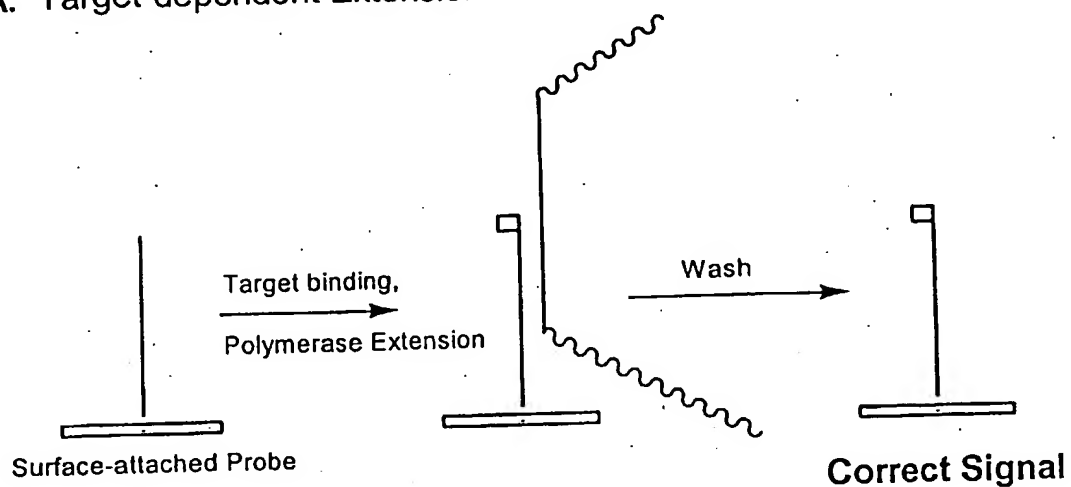
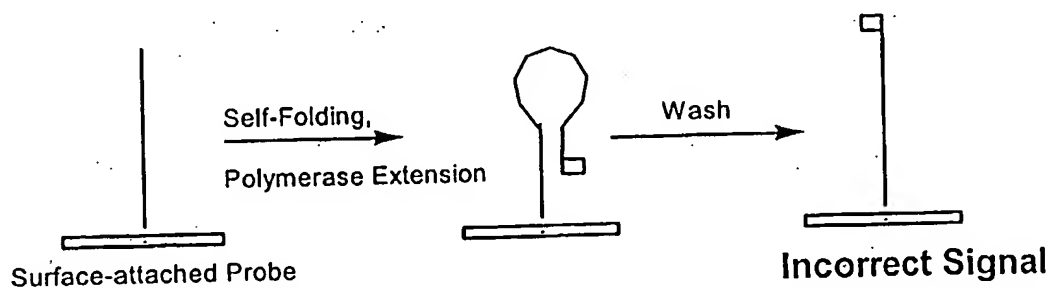


Fig 1B

A. Target-dependent Extension**B. Target-independent Extension**

450 BNP CHINA

SNP ID	GENE	POSITION	FROM TRANSLATION START	SEQ. SOURCE	EXON/INTRON LOCATION	MUTATION	CONSEQUENCE	ENZYME EFFECT (IN YMD)
208 1618	CYP 2D6	1618	-1	GB M333081	5' FLANK		N/A	
208 1620	CYP 2D6	1620	10	GB M333186	EXON 1	G>A	V71M	
208 1623	CYP 2D6	1623	31	GB M333369	EXON 1	G>A	V11M	
208 1626	CYP 2D6	1626	77	GB M333369	EXON 1	G>T	R20H	
208 1696	CYP 2D6	1696	82	GB M333369	EXON 1	G>T	P21C	
208 1701	CYP 2D6	1701	100	GB M333369	EXON 1	G>T	P21C	
208 1711	CYP 2D6	1711	124	GB M333369	EXON 1	G>A	G42R	NO ACTIVITY
208 1743	CYP 2D6	1743	128	GB M333369	EXON 1	G>T	FRAMESHIFT	
208 1757	CYP 2D6	1757	138	GB M333369	INTRON 1	G>C	SPICE DEFECT	
208 2202	CYP 2D6	2202	2562	GB M333369	EXON 2	G>T	A 85 V	
208 2276	CYP 2D6	2276	2578	GB M333369	EXON 2	G>A	L 91 M	
208 2503	CYP 2D6	2503	2578	GB M333369	EXON 2	A>G	H 64 R	
208 2516	CYP 2D6	2516	2603	GB M333369	EXON 2	D>G	SILENT	
208 2517	CYP 2D6	2517	2612	GB M333369	EXON 2	G>T	T 107 I	
208 2542	CYP 2D6	2542	2642	GB M333369	EXON 2	G>T	V 158 M	
208 2553	CYP 2D6	2553	1559	GB M333369	EXON 3	G>A	SILENT	
208 2571	CYP 2D6	2571	1659	GB M333369	EXON 3	G>C	O 151 E	NO ACTIVITY
208 2590	CYP 2D6	2590	1704	GB M333369	EXON 3	G>G	FRAMESHIFT	
208 3225	CYP 2D6	3225	1707	GB M333369	EXON 3	T>G	SILENT	
208 3376	CYP 2D6	3376	1724	GB M333369	EXON 3	A>G	N 188 D	
208 3343	CYP 2D6	3343	1740	GB M333369	EXON 3	A>G	G 164 GTP	
208 3368	CYP 2D6	3368	1768	GB M333369	EXON 3	G>T	SPICE DEFECT	
208 3477	CYP 2D6	3477	1768	GB M333369	EXON 4	G>A	R 143 C	
208 3485	CYP 2D6	3485	1856	GB M333369	EXON 4	T>C	G>A	
208 3488	CYP 2D6	3488	1869	GB M333369	EXON 4	G>A	R 201 H	
208 3522	CYP 2D6	3522	1843	GB M333369	EXON 4	H>G	FRAMESHIFT	
208 3602	CYP 2D6	3602	1978	GB M333369	EXON 4	G>A	G 212 E	NORMAL
208 3603	CYP 2D6	3603	1978	GB M333369	EXON 4	G>T	SILENT	
208 3695	CYP 2D6	3695	1978	GB M333369	EXON 4	T>C	SILENT	
208 3797	CYP 2D6	3797	1978	GB M333369	EXON 4	T>C	SILENT	
208 3838	CYP 2D6	3838	2170	GB M333369	EXON 5	G>T	A 271 S	NORMAL
208 4059	CYP 2D6	4059	2460	GB M333369	EXON 5	G>T	FRAMESHIFT	NO ACTIVITY
208 4102	CYP 2D6	4102	2483	GB M333369	EXON 5	A>G	FRAMESHIFT	
208 4188	CYP 2D6	4188	2575	GB M333369	EXON 5	G>A	FRAMESHIFT	NO ACTIVITY
208 4194	CYP 2D6	4194	2587	GB M333369	EXON 5	G>A	FRAMESHIFT	NO ACTIVITY
208 4206	CYP 2D6	4206	2613	GB M333369	EXON 5	G>A	FRAMESHIFT	NO ACTIVITY
208 4222	CYP 2D6	4222	2613	GB M333369	EXON 5	G>A	FRAMESHIFT	NO ACTIVITY
208 4263	CYP 2D6	4263	2653	GB M333369	EXON 5	G>T	R 201 C	
208 4272	CYP 2D6	4272	2653	GB M333369	EXON 5	A>C	L 207 L	NONE
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208 4365	CYP 2D6	4365	2653	GB M333369	EXON 5	G>C	E 419 Q	
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208 4446	CYP 2D6	4446	2653	GB M333369	EXON 5	G>C	G>C	
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208 4448	CYP 2D6	4448	2653	GB M333369	EXON 5	G>C	G>C	
208 4449	CYP 2D6	4449	2653	GB M333369	EXON 5	G>C	G>C	
208 4450	CYP 2D6	4450	2653	GB M333369	EXON 5	G>C	G>C	
208 4451	CYP 2D6	4451	2653	GB M333369	EXON 5	G>C	G>C	
208 4452	CYP 2D6	4452	2653	GB M333369	EXON 5	G>C	G>C	
208 4453	CYP 2D6	4453	2653	GB M333369	EXON 5	G>C	G>C	
208 4454	CYP 2D6	4454	2653	GB M333369	EXON 5	G>C	G>C	
208 4455	CYP 2D6	4455	2653	GB M333369	EXON 5	G>C	G>C	
208 4456	CYP 2D6	4456	2653	GB M333369	EXON 5	G>C	G>C	
208 4457	CYP 2D6	4457	2653	GB M333369	EXON			

Fig. 3D

ALLELE/HAPLOTYPE FAM	IUPAC CODE	ALLELE 1	ALLELE 2	SNP SOURCE INFO
	R	G	A	Falbrother, RS: Pharmacogenetics 1998
'5A'	W	T	A	Parsons, L: FEBS Lett 1993
	S	C	G	McBride, ON: Molec Adv Res 1997
'3	R	G	A	Hu, Y: Mol Pharmacology 1997
	Y	C	T	GeneBank SNP database, [D50111]
	R	A	G	HGBase, [SNP000000037]
	R	A	G	OMIM, [124410.0001]
	R	A	G	Smart and Dally: Pharmacogenetics 1999
	R	G	A	Smart and Dally: Pharmacogenetics 1999
	Y	C	T	Corcoran, L: Cancer Res 1998
'4	M	C	A	Hayashi, S: J. Biochem 1991
'2B', '2C'	R	A	G	HGBase, [SNP000000286]
'3	Y	T	C	GeneBank SNP database, [D12572]
	Y	T	C	Stollow, L: Am. J. Human Genetics 1998
'2	S	T	C	Stollow, L: Am. J. Human Genetics 1998
'11	S	G	C	Stollow, L: Am. J. Human Genetics 1998
'12	R	G	A	Stollow, L: Am. J. Human Genetics 1998
	Y	T	C	Bejjani Hum Mol Genetics 2000
'2	K	G	T	Stollow, L: Am. J. Human Genetics 1998
'13	K	G	T	Stollow, L: Am. J. Human Genetics 1998
'14	K	G	T	Stollow, L: Am. J. Human Genetics 1998
'15	K	G	T	Stollow, L: Am. J. Human Genetics 1998
'18	R	G	A	Bejjani: Hum Mol Genetics 2000
	R	G	A	Bejjani: Am J Hum Genetics 1998
'19	R	G	A	Stollow, L: Am. J. Human Genetics 1998
'20	Y	C	T	Stollow, L: Am. J. Human Genetics 1998
	R	G	A	Stollow, L: Am. J. Human Genetics 1998
'3	S	C	G	Stollow, L: Am. J. Human Genetics 1998
	Y	C	T	Stollow, L: Am. J. Human Genetics 1998
'4	Y	T	C	Stollow, L: Am. J. Human Genetics 1998
'25	R	A	G	Stollow, L: Am. J. Human Genetics 1998
	Y	C	T	Stollow, L: Am. J. Human Genetics 1998
	S	C	G	HGBase, [SNP000000047]
	W	T	A	HGBase, [SNP00001727] NCBI [12672]
	K	T	G	NCBI, [10916]
'1F'	M	C	A	Sachdev, C: Br J Clin Pharmacol 1999
'2	S	C	G	Huang, JD: Drug Metab Dispos 1999
'2B'	Y	C	T	Richardson, TH: Arch Biochem Biophys 1995
'8	S	G	C	Isaacs, GC: J. Pharmacol Exp Ther 1998
	R	G	A	Boonin, GC: J. Pharmacol Exp Ther 1998
	Y	C	T	HGBase, [SNP000000188]
'3	R	G	A	De Moraes, SM: Mol Pharmacol 1994
'2A	R	G	A	De Moraes, SM: J Biol Chem 1994

CYP2D6 and its pseudogenes: similarity by region
 S. Kimura et al. *Am J Hum Genet* (1989) 45:889-904

Fig 4

	Length, bp			Similarity, %		
	CYP2D6	CYP2D7	CYP2D8	D6/D7	D7/D8	D6/D8
UPSTREAM	774	777	265	97	92	
		186	183			89
	189		186			
EXON 1	268	269	265	97	94	93
INTRON 1	703	701	1620*	98	90	89
EXON 2	172	172	172	95	94	91
INTRON 2	550	528	546	74	78	77
EXON 3	153	153	153	98	93	92
INTRON 3	88	88	88	98	91	93
EXON 4	161	161	161	98	89	91
INTRON 4	433	425	449	94	85	86
EXON 5	177	177	177	99	93	92
INTRON 5	190	192	186	97	84	83
EXON 6	142	142	142	94	92	96
INTRON 6	207	194	204	82	87	90
EXON 7	188	188	185	98	94	95
INTRON 7	454	454	449	98	91	91
EXON 8	142	142	142	99	96	96
INTRON 8	98	98	96	100	97	97
EXON 9	252	252		94		
3'-FLANKING	180	180	181		95	
	538	528	181	97		92

* 3 A/u repeats insertion

50
51
52

GENE	SEQ ID	START POSITION	DIRECTION	SEQ. SOURCE	EXON/INTRON LOCATION	SEQUENCE	PRIMER LENGTH	Ym	BLAST RESULTS
CYP 2D6	1278	FORWARD	GB [M33388]	5' FLANK	CCACAGAGCTTTGAGAGCTTCA	23	71	ACCEPTABLE	
CYP 2D6	5790	REVERSE	GB [M33388]	EXON 9	CTCACAGGGAAGCAAGACAGCAAT	25	67	ACCEPTABLE	
CYP 2E1	1487	FORWARD	GB [J02843]	5' FLANK	ATGAAAGCTCTGCGGAGGAGGCA	22	71	ACCEPTABLE	
CYP 2E1	4022	REVERSE	GB [J02843]	EXON 2	GAGAAATCTCTCTGAGTGTGCA	25	65	ACCEPTABLE	
CYP 2E1	7445	FORWARD	GB [J02843]	INTRON 3	CACCTTCTCAACAGCCTCTGTGCA	24	69	ACCEPTABLE	
CYP 2E1	10659	REVERSE	GB [J02843]	INTRON 6	CACCTGTGCTCTCTGAGGTTTAA	23	68	ACCEPTABLE	
CYP 2E1	12667	FORWARD	GB [J02843]	INTRON 8	ACCTGAGCCCTGACAGCTTCTTACTTAA	27	68	ACCEPTABLE	
CYP 2E1	19330	REVERSE	GB [J02843]	INTRON 8	CCTGAGGCTGTGGCTTGTTCGGATGAA	26	68	ACCEPTABLE	
CYP 3A4	747	FORWARD	GB ID11131	5' FLANK	CACACCACTCAGCTGACCTCTCTTGA	25	67	ACCEPTABLE	
CYP 3A4	841	REVERSE	GB ID11131	5' FLANK	CTCTGCTTAACGGGAGCTATGGGTGAA	27	87	ACCEPTABLE	
CYP 1A1	7080	FORWARD	GB [D04300]	EXON 7	CATCGAGCTCAATGCAAGCTAGATAGAA	30	70	ACCEPTABLE	
CYP 1A1	8206	REVERSE	GB [D04300]	3' FLANK	CTCTCGACGCGAAGGGGACTCA	21	68	ACCEPTABLE	
CYP 1B1	3669	FORWARD	GB [U56435]	INTRON 1	GCAGACAGGATTAAGAGCTGCTCA	25	69	ACCEPTABLE	
CYP 1B1	4704	REVERSE	GB [U56435]	EXON 2	ATCTGTATGTGACAGCTCGAGTGGCA	25	70	ACCEPTABLE	
CYP 1B1	7883	FORWARD	GB [U56435]	EXON 3	CAGACCAAGAGGATACACATCACTTGA	30	70	ACCEPTABLE	
CYP 1B1	8235	REVERSE	GB [U56435]	EXON 3	CCAGCTCTGAGATCTCTGTATGCTCA	27	69	ACCEPTABLE	
CYP 1A2	2995	FORWARD	GB [M31664]	8' FLANK	GGGTCGACATCTCCCGAGGCA	22	74	ACCEPTABLE	
CYP 1A2	2972	REVERSE	GB [M31664]	EXON 1	CTCTGTATCTGCTCTGAGACCACTG	26	70	ACCEPTABLE	
CYP 2C19	197	FORWARD	GB [NM 000769]	EXON 2	CTCTGTATCTGCTCTGAGACCACTG	26	70	ACCEPTABLE	
CYP 2C19	787	REVERSE	GB [NM 000769]	EXON 5	TCCCGAGGCTTGTGTGATGCTATC	24	70	ACCEPTABLE	

PROBE ID	SEQUENCE
CYP1A1 V 2 70.6568.A.S	GCAAGCGGAAGTGTATCGGTGAGAA
CYP1A1 V 2 70.8568.C.S	GCAAGCGGAAGTGTATCGGTGAGAC
CYP1A1 60.6570.A.A	TCCCAGCGGGCAAT
CYP1A1 60.6570.G.A	TCCCAGCGGGCAAC
CYP1A1 V 2 60.7320.C.A	ATAAGGGTCTTACAAGGCCG
CYP1A1 V 2 60.7320.T.A	AATAAGGGTCTTACAAGGCCA
CYP1A2 60+1.2640.A.A	CATCTACCATGCGTCCTGTG
CYP1A2 60+1.2640.C.A	ATCTACCATGCGTCCTGGG
CYP1A2 V2.2868.C.S	TGGCCTCTGCCATCTTCT
CYP1A2 V2.2868.G.S	TGGCCTCTGCCATCTTG
CYP1A2 V3.2866.C.S	TGGCCTCTGCCATCTTCT
CYP1A2 V3.2866.G.S	TGGCCTCTGCCATCTTGT
CYP1B1 60.3793.C.A	CCATGCTGGGGACAGAG
CYP1B1 60.3793.T.A	CCATGCTGGGGACAGAA
CYP1B1 60.3947.C.S	GAGGCGGCAGCTCC
CYP1B1 60.3947.G.S	GAGGCGGCAGCTCG
CYP1B1 60.3976.C.S	GCCCGTTTGCCTGC
CYP1B1 60.3976.G.S	GCCCGTTTGCCTGG
CYP1B1 60.3987.A.A	GCCGCCGCGTTTT
CYP1B1 60.3987.G.A	GCCGCCGCGTTTC
1B1 4035.C.S	CGTTCGCTCGCCC
1B1 4035.T.S	CTCGTTCGCTCGCCT
CYP1B1 60+2-1.4160.G.A	GAAGGAGGCGAAGGCCG
CYP1B1 60+2-1.4180.T.A	GAAGGAGGCGAAGGACG
1B1 V2.4306.A.A	TCAGCACGTGGCCCT
1B1 V2.4306.T.A	CAGCACGTGGCCCAG
CYP1B1 60+1.4646.G.S	AGTTCTTGAGGCACTGCGA
CYP1B1 60+1.4646.T.S	CAAGTTCTTGAGGCACTGCTA
1B1 4668.C.A	TCGCGGGGGGG
1B1 4668.G.A	TCGCGGGGGGC
CYP1B1 60.7930.G.S	GAATTGGATCAGGTCGTGG
CYP1B1 60.7930.T.S	AGAATTGGATCAGGTCGTGT
1B1 V2.7940.A.A	TGGTIACCCATACAAGGCAGAT
1B1 V2.7940.G.A	GGTIACCCATACAAGGCAGACG
CYP1B1 60+1.7857.A.S	CGTCTGCCTTGTATGGGTAA
CYP1B1 60+1.7857.G.S	CGTCTGCCTTGTATGGGTGA
CYP1B1 60.7973.C.A	GGAAGGCCAGGACATAGG
CYP1B1 60.7973.T.A	AGGAAGGCCAGGACATAGA
CYP1B1 60.7996.A.S	TATGTCTCTGGCCTTCCTTTATA
CYP1B1 60.7996.G.S	GTCCTGGCCTTCCTTTATG
CYP1B1 60+1.8131.C.S	GTCTGTGAATCATGACCCACT
CYP1B1 60+1.8131.G.S	GTCTGTGAATCATGACCCAGT
CYP1B1 60+1*.8184.C.A	GTCTTGITGATGAGGCCGT
CYP1B1 60+1*.8184.T.A	GTCCTTGITGATGAGGCCAT
CYP1B1 60.8195.A.A	TGCTGGTCAGGTCCTTGT
CYP1B1 60.8195.G.A	GCTGGTCAGGTCCTTGC
CYP1B1 60.8242.C.S	TTCAAGTGGGCAAAAGGC
CYP1B1 60.8242.T.S	TTTTCAGTGGGCAAAAGGT
CYP1B1 60.8587.C.S	TCAATTAGCGTTTAAGGTGAGC
CYP1B1 60.8587.G.S	TCAATTAGCGTTTAAGGTGAGG
CYP1B1 60.8807.A.S	CCCAAACACTTACACCAAACA
CYP1B1 60.8807.T.S	ACCCAAACACTTACACCAAAC
CYP1B1 60+1.9184.G.S	GAGTATAGTGGGGTTCCATGAGT
CYP1B1 60+1.9184.T.S	GAGTATAGTGGGGTTCCATGATT
CYP2C19EXONS 70.276.C.A	GAAATGGCCTCTTCCAGAAAAC
CYP2C19EXONS 70.276.G.A	GGAAATGGCCTCTTCCAGAAAAC
CYP2C19EXONS 70.395.A.A	CTCCTCTTCCCCATCCCAAATCT
CYP2C19EXONS 70.395.G.A	CCTCTTCCCCATCCCAAATCC

Fig 6/

CYP2C19EXONS 70.430.T.A	AGCGGGCTTCCTCTTGAACACA
CYP2C19EXONS 60.636.A.S	GATTGTAAGCACCCCTGA
CYP2C19EXONS 60.636.G.S	TTGTAAGCACCCCTGG
CYP2C19EXONS 60.681.A.S	CCACTATCATTGATTATTCCCA
CYP2C19EXONS 60.681.G.S	CCACTATCATTGATTATTCCCG
CYP2D6 70.1638.A.S	AGGCAGITATGGGGCTAGAAGCACTGA
CYP2D6 70.1638.G.S	GGCAGITATGGGGCTAGAAGCACTGG
CYP2D6 70.1650.A.A	AGGAGCAGGAAGATGGCCACTATCAT
CYP2D6 70.1650.G.A	GGAGCAGGAAGATGGCCACTATCAC
CYP2D6 70.1698.A.S	GGACCTGATGCACCGGCA
CYP2D6 70.1698.G.S	GGACCTGATGCACCGGCG
CYP2D6 70.1701.C.A	TGIGTAGCGTGCAGCCCAGCG
CYP2D6 70.1701.T.A	GTGIGTAGCGTGCAGCCCAGCA
CYP2D6 60.1719.C.A	GGGGGCCTGGTGG
CYP2D6 60.1719.T.A	AGGGGGCCTGGTGA
CYP2D6 70.1743.A.S	CCCCCTGCCACTGCCCA
CYP2D6 70.1743.G.S	CCCCTGCCACTGCCCG
2D6H.1757.G.S	CCCTGCCACTGCCCIGGCTGGGCAACCTG
2D6H.1757.T.S	CCCTGCCACTGCCCIGGCTGGGCAACCTT
2D6H V2.1757.G.S	CCTGCCACTGCCCHGGCTGGGCAACCTGCT
2D6H V2.1757.T.S	CCTGCCACTGCCCHGGCTGGGCAACCTTCT
CYP2D6 60+1.2502.C.A	CGGCGCCGCAAGT
CYP2D6 60+1.2502.G.A	CGGCGCCGCAACT
CYP2D6 60+1.2502.C.S	TGACCCTCCCTCTGCACT
CYP2D6 60+1.2502.G.S	TGACCCTCCCTCTGCAGT
CYP2D6 60.2578.C.S	GCTCAATGGGCTGGC
CYP2D6 60.2578.T.S	GTGCTCAATGGGCTGGT
CYP2D6 60.2593.A.A	CGCCGIGGGTCACCAT
CYP2D6 60.2593.C.A	CGCCGIGGGTCACCAG
CYP2D6 70+*.2603.A.S	GAGGCGITGGTGACCCACG
CYP2D6 70+*.2603.G.S	CGAGGCGITGGTGACCCG
CYP2D6 60+2-1.2816.C.A	GCGGTGCGCGGT
CYP2D6 60+2-1.2818.G.A	GGCGGTGCGCGGT
CYP2D6 60+1.2842.C.S	GCCTGTGCCCATCACC
CYP2D6 60+1.2842.T.S	CGCCTGTGCCCATCATC
CYP2D6 60+2.2642.C.A	CCIAAACCAGGATCTGGGTG
CYP2D6 60+2.2642.T.A	CCIAAACCAGGATCTGGATG
CYP2D6 70+1.2658.C.A	TGGGAACGCGGCCCGA
CYP2D6 70+1.2658.T.A	TGGGAACGCGGCCCAA
CYP2D6 60+1.3278.A.S	CAGAGGCGCTTCTCCAT
CYP2D6 60+1.3278.G.S	CAGAGGCGCTTCTCCGT
CYP2D6 60.3280.C.S	CAGAGGCGCTTCTCCITC
CYP2D6 60.3280.G.S	CAGAGGCGCTTCTCCITG
CYP2D6 70+1.3323.C.S	TGGGCAAGAAGTCGCTGGAGCA
CYP2D6 70+1.3323.G.S	TGGGCAAGAAGTCGCTGGAGGA
2D6H.3326.G.A	GCIGCCTCCTCGGTACCCC
2D6H.3326.T.A	GCIGCCTCCTCGGTACCCA
2D6H V2.3328.G.A	GCIGCCTCCTCGGTACCCCT
2D6H V2.3326.T.A	GCIGCCTCCTCGGTACCCAC
CYP2D6 70.3343.C.A	CGGCACAAAGGCAGGCG
CYP2D6 70.3343.T.A	GCGGCACAAAGGCAGGCA
CYP2D6 70.3368.A.S	TGTGCCGCTTCGCCA
CYP2D6 70.3388.G.S	GTGCCGCTTCGCCG
CYP2D6 70.3377.G.S	CGCCTTCGCCACCACTCCG
CYP2D6 70.3377.T.S	CCGCTTCGCCACCACTCCT
CYP2D6 60.3465.A.S	CATCTCCCACCCCCAA
CYP2D6 60.3465.G.S	CATCTCCCACCCCCAG
CYP2D6 60.3477.C.A	AGAGICCGTTGGGGCG
CYP2D6 60.3477.T.A	AAGAGICCGTTGGGGCA
CYP2D6 60.3488.C.A	CGGCTTGTCCAAGAGG

Fig 6B

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CYP2D6 60.3582.A.A	CCAGCAGCCTGAGGAAGT
CYP2D6 60.3582.G.A	CAGCAGCCTGAGGAAGC
2D6H.3592.A.S	TCAGGCTGCTGGACCTAGCTCAGGA
2D6H.3592.G.S	CAGGCTGCTGGACCTAGCTCAGGGA
CYP2D6 60.3595.A.S	TGGACCTAGCTCAGGAGGA
CYP2D6 60.3595.G.S	GGACCTAGCTCAGGAGGG
CYP2D6 70.3597.C.S	GCTGCTGGACCTAGCTCAGGAGGIAC
CYP2D6 70.3597.T.S	GCTGCTGGACCTAGCTCAGGAGGIAT
CYP2D6 60.3598.C.A	CCCGACTCCTCCTTCG
CYP2D6 60.3598.T.A	GCCCGACTCCTCCTTCA
CYP2D6 70.4099.C.S	TCCTCCTGCAIATCCCAGCGC
CYP2D6 70.4099.T.S	GTCTCCTGCAIATCCCAGCGT
2D6H V2.4168.A.S	CTGGATGAGCTGCTAACTGAGCACAGG
2D6H V2.4168.G.S	CTGGATGAGCTGCTAACTGAGCACGGG
2D6H V3.4168.A.S	GCTGGATGAGCTGCTAACTGAGCACACA
2D6H V3.4168.G.S	GCTGGATGAGCTGCTAACTGAGCACGG
CYP2D6 70.4194.A.S	GGGACCCAGCCCAGCCA
CYP2D6 70.4194.C.S	GGGACCCAGCCCAGCCC
2D6H.4206.G.S	GCCCAGCCICCCGAGACCTGAGG
2D6H.4208.T.S	GCCCAGCCICCCGAGACCTGACT
2D6H.4232.A.A	TGGCAGCCACTCTCACCTTCT
2D6H.4232.G.A	TGGCAGCCACTCTCACCTC
CYP2D6 60.4489.C.S	GCTTCAATGATGAGAACCTGC
CYP2D6 60.4489.T.S	AGCTTCAATGATGAGAACCTGT
CYP2D6 70.4472.A.A	GCAGAGAACAGGTCAGCCACCACTAT
CYP2D6 70.4472.C.A	GCAGAGAACAGGTCAGCCACCACTAG
CYP2D6 V3.4554.C.A	TGGGCTCACGCTGCACATCIIGAGG
CYP2D6 V3.4554.A.A	GCTCACGCTGCACATCIIGAT
CYP2D6 V2.4554.C.A	GCTCACGCTGCACATCIIGAGG
CYP2D6 V2.4554.A.A	GCTCACGCTGCACATCIIGAT
CYP2D6 70.4557.C.S	GGCCTCCTGCTCATGATCCTACITCC
CYP2D6 70.4557.T.S	GGCCTCCTGCTCATGATCCTACITCT
CYP2D6 70.4558.A.A	TGGGCTCACGCTGCACATCT
CYP2D6 70.4558.G.A	GGGCTCACGCTGCACATCC
CYP2D6 70.4802.A.S	GTGTCCAACAGGAGATCGACGACA
CYP2D6 70.4802.G.S	TGTCCAACAGGAGATCGACGACG
CYP2D6 70+*.4817.C.S	TCGACGACITGATAGGGCAGGTGCGG
CYP2D6 70+*.4817.G.S	ATCGACGACITGATAGGGCAGGTGGG
CYP2D6 70.4896.C.S	TGCAGCGCTTTGGGGACAC
CYP2D6 70.4896.T.S	GTGCAGCGCTTTGGGGACAT
CYP2D6 60.4907.A.S	GGACAICGTCCCCCTGA
CYP2D6 60.4907.G.S	GGACAICGTCCCCCTGG
CYP2D6 70.5447.A.A	AGACGGCCTCATCCTTCAGCACT
CYP2D6 70.5447.G.A	ACGGCCTCATCCTTCAGCACC
CYP2D6 70.5472.A.S	CTGAAGGATGAGGCCGTCTGGA
CYP2D6 70.5472.G.S	TGAAGGATGAGGCCGTCTGGG
CYP2D6 70.5496.C.S	CCTTCCGCTTCCACCCCC
CYP2D6 70.5496.G.S	CCTTCCGCTTCCACCCCCG
CYP2D6 70+1.5508.C.S	CGCTTCCACCCCCAACACTTCCCCG
CYP2D6 70+1.5506.T.S	CCGCTTCCACCCCCAACACTTCCCTG
CYP2D6 70+1.5661.A.S	CCCCTCCCCACAGGCCAC
CYP2D6 70+1.5661.G.S	CCCTCCCCACAGGCCGC
CYP2D6 70.5734.C.A	CAGTGGGCACCGAGAAGCTG
CYP2D6 70.5734.T.A	TCCAGTGGGCACCGAGAAGCTA
CYP2E1 60-1+1.1532.C.A	CTGCACCTAACACTGCAGC
CYP2E1 60-1+1.1532.G.A	CTGCACCTAACACTGCACC
CYP2E1 60.1627.C.A	CATTCTATACITGTATTTATACAAAAATGAGAG
CYP2E1 60.1627.G.A	CATTCTATACITGTATTTATACAAAAATGAGAC
CYP2E1 V2.1772.C.A	TCTTAATTCATAGGTTGCAATTTTGT
CYP2E1 V2.1772.T.A	TTCTTAATTCATAGGTTGCAATTTTATA

Fig 6C

CYP2E1_V3.1800.T.S	TTGCAACCTATGAATTAAGAACTTCTA
CYP2E1_V3.1800.C.S	ATTGCAACCTATGAATTAAGAACTCC
CYP2E1_60+1.2019.C.A	GATTTGTTTTACATTAGGGTAAATTTGG
CYP2E1_60+1.2019.T.A	GGATTTGTTTTACATTAGGGTAAATTTAG
2E1_2492.A.A	GTGGGGTGAGGTACCGT
2E1_2492.T.A	GTGGGGTGAGGTACCGA
2E1_2492.A.S	TGCCAAAGGGCAGGA
2E1_2492.T.S	GTGCCAAAGGGCAGGT
2E1_2473.A.A	GCCCTTTGGCACTGGT
2E1_2473.G.A	CCCTTTGGCACTGGC
2E1_2473.A.S	GGAGTCCCCGTTGTCTAA
2E1_2473.G.S	GGAGTCCCCGTTGTCTAG
CYP2E1_60.2754.G.S	GGGTCACCCTCCTTCTCAG
CYP2E1_60.2754.T.S	GGGTCACCCTCCTTCTCAT
CYP2E1_60+2.3958.A.S	GTGGGCTCGCAGCACA
CYP2E1_60+2.3958.G.S	TGGGCTCGCAGCGCA
CYP2E1_V2.3858.A.A	CCGTGCATCACCACCATGT
CYP2E1_V2.3858.G.A	GTGCATCACCACCATGOG
CYP2E1_60.10458.A.S	CACACCCAGCTGATTAAAAATTA
CYP2E1_60.10458.T.S	CACACCCAGCTGATTAAAAATTT
CYP2E1_60.12720.C.S	TCACTAAGCAACTCCTTCAACTC
CYP2E1_60.12720.G.S	TCACTAAGCAACTCCTTCAACTG
CYP2E1_60.12847.A.S	TTTCTCCTAGGGCACAGTCA
CYP2E1_60.12847.G.S	TCTCCTAGGGCACAGTCG
CYP2E1_60.12945.C.A	GGCTTGAAATAGTCACTGTACTTG
CYP2E1_60.12945.T.A	AATGGCTTGAAATAGTCACTGTACTTA
CYP3A4_60.816.A.S	GCCATAGAGACAAGGGCAA
CYP3A4_60.818.G.S	GCCATAGAGACAAGGGCAG
CYP3A4_60.818.A.S	CCAGTAACATTGATTGAGTTGTTTA
CYP3A4_60.818.G.S	CAGTAACATTGATTGAGTTGTTTG
Amplicon Control Probes	
1A1.23F22R_A.X.A	GCAGGATCCCTTAGGCTTG
1A1.23F22R_B.X.S	AGCCAGGAGGCCTGCTA
1A2.5F3R_A.X.S	TATCCAGCTGGGAGCCAA
1A2.5F3R_B.X.S	CCAGCCCCATGGCTCT
1B1.2F4R_A.X.S	CACGACGACCCGAGTT
1B1.2F4R_B.X.S	CGGTGCGCACCGTT
1B1.8F11R_A.X.A	TTGGGTTGGCCCTGAA
1B1.8F11R_B.X.S	TGGGCTATGCAGGAGCTT
2C19.3F6R_A.X.A	GCACAGCCCAGGATGAA
2C19.3F6R_B.X.A	CATGCAGCACCACCATG
2D6.1F1R_A.X.S	AGCCCATTTGGTAGTGAGGCAGG
2D6.1F1R_B.X.S	GAGCCCATTTGGTAGTGAGGCAGA
2E1.1F6R_A.X.A	AGGITGGTATTGAACAACCACAA
2E1.1F6R_B.X.A	ATTGAGGTAATTCACAACAGGC
2E1.8F19R_A.X.S	GACTGTGGCCGACCTGTT
2E1.8F19R_B.X.S	GCACAGTGCAGAGCGCTT
2E1.11F13R_A.X.S	CCAGATGAAAGCCACATT
2E1.11F13R_B.X.S	AAGCCACATTTTGTTAACATG
3A4.1F1R_A.X.S	GCTTGTTGGGATGAATTTCAA
3A4.1F1R_B.X.S	CTGATAAGAACCCAGAACCCTT

Fig 6D

File 7

1A1	1A2				Comments
NONE	CYP 1A1	1213	CYP1A1 V 2 60.1213.A.S	CYP1A1 V 2 60.1213.G.S	No Amplicon Coverage
NONE	CYP 1A1	1223	CYP1A1 V 2 60.1223.C.A	CYP1A1 V 2 60.1223.T.A	No Amplicon Coverage
23F22R	CYP 1A1	6568	CYP1A1 V 2 70.6568.A.S	CYP1A1 V 2 70.6568.C.S	
23F22R	CYP 1A1	6570	CYP1A1 60.6570.A.A	CYP1A1 60.6570.G.A	
23F22R	CYP 1A1		CYP1A1 V 2 70.6570.A.A	CYP1A1 V 2 70.6570.G.A	Remove unneeded redundancy
NONE	CYP 1A1	7320	CYP1A1 V 2 60.7320.C.A	CYP1A1 V 2 60.7320.T.A	
23F22R	CYP 1A1	7547	CYP1A1 60+1.7547.A.S	CYP1A1 60+1.7547.T.S	No Amplicon Coverage
23F22R	CYP 1A1	CNTRL	1A1.23F22R A.X.A		Independent Amplicon Controls
23F22R	CYP 1A1	CNTRL	1A1.23F22R B.X.S		Independent Amplicon Controls
5F3R	CYP 1A2	2640	CYP1A2 60+1.2640.A.A	CYP1A2 60+1.2640.C.A	Redesign to overcome possible selfX
5F3R	CYP 1A2		CYP1A2 60+1.2640.A.A	CYP1A2 60+1.2640.C.A	Remove unneeded redundancy
5F3R	CYP 1A2	2866	CYP1A2 V2.2866.C.S	CYP1A2 V2.2866.G.S	Redesign of one Probe (CYP1A2 V2.2866.G.S = CYP1A2 60.2866.G.S)
5F3R	CYP 1A2		CYP1A2 V3.2866.C.S	CYP1A2 V3.2866.G.S	
5F3R	CYP 1A2	CNTRL	1A2.5F3R A.X.S		Independent Amplicon Controls
5F3R	CYP 1A2	CNTRL	1A2.5F3R B.X.S		Independent Amplicon Controls
2F4R	CYP 1B1	3793	CYP1B1 60.3793.C.A	CYP1B1 60.3793.T.A	
2F4R	CYP 1B1	3947	CYP1B1 60.3947.C.S	CYP1B1 60.3947.G.S	
2F4R	CYP 1B1	3976	CYP1B1 60.3976.C.S	CYP1B1 60.3976.G.S	
2F4R	CYP 1B1	3987	CYP1B1 60.3987.A.A	CYP1B1 60.3987.G.A	
2F4R	CYP 1B1	4035	1B1 4035.C.S	1B1 4035.T.S	Newly Identified SNP
2F4R	CYP 1B1	4160	CYP1B1 60+2-1.4160.G.A	CYP1B1 60+2-1.4160.T.A	Redesigned to overcome GG self extension
2F4R	CYP 1B1	4308	1B1 V2.4308.A.A	1B1 V2.4308.T.A	Newly Identified SNP
2F4R	CYP 1B1	4646	CYP1B1 60+1.4646.G.S	CYP1B1 60+1.4646.T.S	
2F4R	CYP 1B1	4668	1B1 4668.C.A	1B1 4668.G.A	Newly Identified SNP
2F4R	CYP 1B1	CNTRL	1B1.2F4R A.X.S		Independent Amplicon Controls
2F4R	CYP 1B1	CNTRL	1B1.2F4R B.X.S		Independent Amplicon Controls

SNP ID	PROBE ID
2E1_1772	CYP2E1_60.1772.C.A
2E1_1772	CYP2E1_60.1772.T.A
2E1_2019	CYP2E1_60.2019.C.S
2E1_2019	CYP2E1_60.2019.T.S
2E1_2754	CYP2E1_60.2754.G.S
2E1_2754	CYP2E1_60.2754.T.S
2E1_3085	CYP2E1_60.3085.A.A
2E1_3085	CYP2E1_60.3085.T.A
2E1_3104	CYP2E1_60.3104.A.A
2E1_3104	CYP2E1_60.3104.G.A
2E1_7592	CYP2E1_60.7592.A.S
2E1_7592	CYP2E1_60.7592.G.S
2E1_10456	CYP2E1_60.10456.A.S
2E1_10456	CYP2E1_60.10456.T.S
2E1_12720	CYP2E1_60.12720.C.S
2E1_12720	CYP2E1_60.12720.G.S
2E1_12847	CYP2E1_60.12847.A.S
2E1_12847	CYP2E1_60.12847.G.S
2E1_12945	CYP2E1_60.12945.C.A
2E1_12945	CYP2E1_60.12945.T.A
2D6_4802	CYP2D6_70.4802.A.S
2D6_4802	CYP2D6_70.4802.G.S
2D6_4896	CYP2D6_70.4896.C.S
2D6_4896	CYP2D6_70.4896.T.S
2D6_4907	CYP2D6_70.4907.A.S
2D6_4907	CYP2D6_70.4907.G.S
2D6_4907	CYP2D6_70.4907.A.S
2D6_4907	CYP2D6_70.4907.G.S
2D6_5447	CYP2D6_70.5447.A.A
2D6_5447	CYP2D6_70.5447.G.A
2D6_5472	CYP2D6_70.5472.A.S
2D6_5472	CYP2D6_70.5472.G.S
2D6_3595	CYP2D6_60.3595.A.A
2D6_3595	CYP2D6_60.3595.G.A
2D6_3595	CYP2D6_60.3595.A.S
2D6_3595	CYP2D6_60.3595.G.S
2D6_3597	CYP2D6_70.3597.C.S
2D6_3597	CYP2D6_70.3597.T.S
2D6_3598	CYP2D6_60.3598.C.A
2D6_3598	CYP2D6_60.3598.T.A
2D6_4089	CYP2D6_70.4089.C.S
2D6_4089	CYP2D6_70.4089.T.S
2D6_4099	CYP2D6_70.4099.C.S
2D6_4099	CYP2D6_70.4099.T.S
2D6_4102	CYP2D6_70.4102.G.A
2D6_4102	CYP2D6_70.4102.T.A
2D6_2642	CYP2D6_60.2642.C.S
2D6_2642	CYP2D6_60.2642.T.S
2D6_2642	CYP2D6_70.2642.C.S
2D6_2642	CYP2D6_70.2642.T.S
2D6_2658	CYP2D6_70.2658.C.A

Fig 8A

2D6 2658	CYP2D6 70.2658.T.A
2D6 3278	CYP2D6 60.3278.A.S
2D6 3278	CYP2D6 60.3278.G.S
2D6 3280	CYP2D6 60.3280.C.S
2D6 3280	CYP2D6 60.3280.G.S
2D6 3343	CYP2D6 70.3343.C.A
2D6 3343	CYP2D6 70.3343.T.A
2D6 1618	CYP2D6 70.1618.A.S
2D6 1618	CYP2D6 70.1618.G.S
2D6 1638	CYP2D6 70.1638.A.S
2D6 1638	CYP2D6 70.1638.G.S
2D6 1650	CYP2D6 70.1650.A.A
2D6 1650	CYP2D6 70.1650.G.A
2D6 1696	CYP2D6 70.1696.A.S
2D6 1696	CYP2D6 70.1696.G.S
2D6 1701	CYP2D6 70.1701.T.A
2D6 1701	CYP2D6 70.1701.C.A
2D6 1719	CYP2D6 60.1719.C.A
2D6 1719	CYP2D6 60.1719.T.A
2D6 1743	CYP2D6 70.1743.A.S
2D6 1743	CYP2D6 70.1743.G.S
2D6 2502	CYP2D6 60.2502.C.A
2D6 2502	CYP2D6 60.2502.G.A
2D6 2578	CYP2D6 60.2578.T.S
2D6 2576	CYP2D6 60.2576.C.S
2D6 2593	CYP2D6 60.2593.A.A
2D6 2593	CYP2D6 60.2593.C.A
2D6 2603	CYP2D6 70.2603.A.S
2D6 2603	CYP2D6 70.2603.G.S
2D6 2616	CYP2D6 60.2616.C.A
2D6 2616	CYP2D6 60.2616.G.A
2D6 3368	CYP2D6 70.3368.A.S
2D6 3368	CYP2D6 70.3368.G.S
2D6 3377	CYP2D6 70.3377.G.S
2D6 3377	CYP2D6 70.3377.T.S
2D6 3485	CYP2D6 60.3465.A.A
2D6 3465	CYP2D6 60.3465.G.A
2D6 3465	CYP2D6 60.3465.A.S
2D6 3465	CYP2D6 60.3465.G.S
2D6 3465	CYP2D6 70.3465.A.A
2D6 3465	CYP2D6 70.3465.G.A
2D6 3485	CYP2D6 70.3465.A.S
2D6 3465	CYP2D6 70.3465.G.S
2D6 3477	CYP2D6 60.3477.C.A
2D6 3477	CYP2D6 60.3477.T.A
2D6 3488	CYP2D6 60.3488.C.A
2D6 3488	CYP2D6 60.3488.T.A
2D6 3562	CYP2D6 60.3562.A.A
2D6 3562	CYP2D6 60.3562.G.A
2D6 4194	CYP2D6 70.4194.A.S
2D6 4194	CYP2D6 70.4194.C.S
2D6 4469	CYP2D6 60.4469.C.S

Fig 8B

2D6 4469	CYP2D6 60.4469.T.S
2D6 4472	CYP2D6 70.4472.A.A
2D6 4472	CYP2D6 70.4472.C.A
2D6 4472	CYP2D6 70.4472.A.S
2D6 4472	CYP2D6 70.4472.C.S
2D6 4554	CYP2D6 70.4554.A.A
2D6 4554	CYP2D6 70.4554.C.A
2D6 4557	CYP2D6 70.4557.C.S
2D6 4557	CYP2D6 70.4557.T.S
2D6 4558	CYP2D6 70.4558.A.A
2D6 4558	CYP2D6 70.4558.G.A
2D6 5496	CYP2D6 70.5496.C.S
2D6 5496	CYP2D6 70.5496.G.S
2D6 5506	CYP2D6 60.5506.C.A
2D6 5506	CYP2D6 60.5506.T.A
2D6 5661	CYP2D6 70.5661.A.S
2D6 5661	CYP2D6 70.5661.G.S
2D6 5734	CYP2D6 70.5734.C.A
2D6 5734	CYP2D6 70.5734.T.A
2D6 5799	CYP2D6 60.5799.C.S
2D6 5799	CYP2D6 60.5799.G.S
2E1 1627	CYP2E1 60.1627.C.A
2E1 1627	CYP2E1 60.1627.G.A
3A4 816	CYP3A4 60.816.G.S
3A4 816	CYP3A4 60.816.A.S
3A4 918	CYP3A4 60.918.A.S
3A4 918	CYP3A4 60.918.G.S
2C19 430	CYP2C19EXONS 70.430.C.A
2C19 430	CYP2C19EXONS 70.430.T.A
2C19 636	CYP2C19EXONS 60.636.A.S
2C19 636	CYP2C19EXONS 60.636.G.S
2C19 681	CYP2C19EXONS 60.681.A.S
2C19 681	CYP2C19EXONS 60.681.G.S
1B1 4160	CYP1B1 60.4160.G.A
1B1 4160	CYP1B1 60.4160.T.A
1B1 7973	CYP1B1 60.7973.C.A
1B1 7973	CYP1B1 60.7973.T.A
1B1 7996	CYP1B1 60.7996.A.S
1B1 7996	CYP1B1 60.7996.G.S
1B1 8006	CYP1B1 60.8006.A.S
1B1 8006	CYP1B1 60.8006.G.S
1B1 8195	CYP1B1 60.8195.A.A
1B1 8195	CYP1B1 60.8195.G.A
1B1 8242	CYP1B1 60.8242.C.S
1B1 8242	CYP1B1 60.8242.T.S
1B1 8587	CYP1B1 60.8587.C.S
1B1 8587	CYP1B1 60.8587.G.S
1A1 1223	CYP1A1 60.1223.C.A
1A1 1223	CYP1A1 60.1223.T.A
2D6 3326	2D6 66.3326.G.A
2D6 3326	2D6 66.3326.T.A
2D6 3326	2D6 66.3326.G.S

Fig 8C

2D6_3326	2D6_66.3326.T.S
2D6_4168	2D6_66.4168.A.A
2D6_4168	2D6_66.4168.C.A
2D6_4168	2D6_66.4168.A.S
2D6_4168	2D6_66.4168.C.S
1A1_6568	CYP1A1_60.6568.A.A
1A1_6568	CYP1A1_60.6568.C.A
1A1_6568	CYP1A1_60.6568.A.S
1A1_6568	CYP1A1_60.6568.C.S
1A1_6570	CYP1A1_60.6570.A.A
1A1_6570	CYP1A1_60.6570.G.A
1A2_2640	CYP1A2_60.2640.A.A
1A2_2640	CYP1A2_60.2640.C.A
1A2_2866	CYP1A2_60.2866.C.S
1A2_2866	CYP1A2_60.2866.G.S
1B1_3793	CYP1B1_60.3793.C.A
1B1_3793	CYP1B1_60.3793.T.A
1B1_3793	CYP1B1_60.3793.C.S
1B1_3793	CYP1B1_60.3793.T.S
1B1_3947	CYP1B1_60.3947.C.S
1B1_3947	CYP1B1_60.3947.G.S
1B1_3976	CYP1B1_60.3976.C.S
1B1_3976	CYP1B1_60.3976.G.S
1B1_3987	CYP1B1_60.3987.A.A
1B1_3987	CYP1B1_60.3987.G.A
1B1_8807	CYP1B1_60.8807.A.S
1B1_8807	CYP1B1_60.8807.T.S
2C19_276	CYP2C19EXONS_70.276.C.A
2C19_276	CYP2C19EXONS_70.276.G.A
2C19_395	CYP2C19EXONS_70.395.A.A
2C19_395	CYP2C19EXONS_70.395.G.A
CONTROL PROBES	
	PBR322WSNPS.4058.C.S
	PBR322WSNPS.4058.T.S
	WIAF-1648.107.A.A
	WIAF-1648.107.G.A
	WIAF-198.38.C.A
	WIAF-198.38.T.A
	2D7PSEUDOGENECONTROL_60.111.C.A
	2D7PSEUDOGENECONTROL_60.111.G.A
	2D7PSEUDOGENECONTROL_60.111.C.S
	2D7PSEUDOGENECONTROL_60.111.G.S
	2D7APSEUDOGENECONTROL_60.23.A.A
	2D7APSEUDOGENECONTROL_60.23.G.A
	2D7APSEUDOGENECONTROL_60.2370.A.S
	2D7APSEUDOGENECONTROL_60.2370.G.S
	2D7APSEUDOGENECONTROL_60.2692.C.S
	2D7APSEUDOGENECONTROL_60.2692.T.S
	2D7APSEUDOGENECONTROL_60.3471.C.A
	2D7APSEUDOGENECONTROL_60.3471.T.A
	2D7PSEUDOGENECONTROL_60.600.G.S

Fig 8D

	2D7PSEUDOGENECONTROL 60.800.T.S
	2D7PSEUDOGENECONTROL 60.1760.C.S
	2D7PSEUDOGENECONTROL 60.1760.T.S
	2D7PSEUDOGENECONTROL 60.2108.A.S
	2D7PSEUDOGENECONTROL 60.2108.G.S
	2D7B 60.3539.G.S
	2D7B 60.3539.T.S
	2D7B 60.3647.A.A
	2D7B 60.3647.G.A
	2D7B 60.3766.A.S
	2D7B 60.3766.C.S
	2D7B 60.4506.C.S
	2D7B 60.4506.G.S
	2D8 60.105.A.A
	2D8 60.105.G.A
	2D8 60.3080.A.S
	2D8 60.3080.G.S
	2D7PSEUDOGENECONTROL 60.1360.C.S
	2D7PSEUDOGENECONTROL 60.1360.T.S
	2D7PSEUDOGENECONTROL 60.3030.A.S
	2D7PSEUDOGENECONTROL 60.3030.G.S
	2D7PSEUDOGENECONTROL 60.3148.A.S
	2D7PSEUDOGENECONTROL 60.3148.G.S
	2D7B 60.442.C.A
	2D7B 60.442.T.A
	2D7B 60.652.G.S
	2D7B 60.652.T.S
	2D7B 60.1185.G.S
	2D7B 60.1185.T.S
	2D7B 60.1316.A.A
	2D7B 60.1316.C.A
	2D7B 60.1671.A.A
	2D7B 60.1671.T.A
	2D7B 60.3172.C.S
	2D7B 60.3172.G.S
	2D8 60.3181.A.A
	2D8 60.3181.G.A
	2D8 60.4120.A.A
	2D8 60.4120.G.A
	2D8 60.4199.C.A
	2D8 60.4199.T.A
	2D8 60.4223.C.S
	2D8 60.4223.T.S
	2D8 60.4750.C.A
	2D8 60.4750.G.A

Fig 8E

Fig 9A

GENE	RELATED/MAPPED TOGETHER	DNA/mRNA/Parlals	ACCESSION #
2D6	2D6	gDNA	M33388
	2D7	gDNA	M33387
	2D7A	gDNA	X58487
	2D7B	gDNA	X58468
	2D8	gDNA	M33387
2E1	2E1	gDNA	J02843
3A4	3A4	Partial gDNA	D11131 AF209389
3A5	3A5	gDNA	AC005020
	3A5P	mRNA Splice Variant (mRNA)	NM_000777 L26985
1A1	1A1	gDNA	X04300
		gDNA	X02612
		Partial	D12525
1A2	1A2	Partial gDNA	M31664
		Partial gDNA	M31665
		Partial gDNA	M31666
		Partial gDNA	M31667
1B1	1B1	gDNA	U56438
2C9	2C9	mRNA	M61855
	2C9	Partial gDNA	L16877
	2C9	Partial gDNA	L16878
	2C9	Partial gDNA	L16879
	2C9	Partial gDNA	L16880
	2C9	Partial gDNA	L16881
	2C9	Partial gDNA	L16882
	2C9	Partial gDNA	L16883

Fig 10A

AMPLICON	LENGTH	SNP ID
1A1 23F 22R	1127	1A1 6568
		1A1 6570
		1A1 7320
1A2 05F 03R	378	1A2 2640
		1A2 2866
1B1 02F 04R	1007	1B1 3793
		1B1 3947
		1B1 3976
		1B1 3987
		1B1 4160
		1B1 4646
1B1 08F 11R	1353	1B1 7930
		1B1 7957
		1B1 7973
		1B1 7996
		1B1 8006
		1B1 8131
		1B1 8147
		1B1 8184
		1B1 8195
		1B1 8242
		1B1 8587
		1B1 8807
		1B1 9164
2C19 03F 06R	-6500	2C19 276
		2C19 395
		2C19 430
		2C19 636
		2C19 681
2D6 01F 01R	4522	2D6 1618
		2D6 1638
		2D6 1650
		2D6 1696
		2D6 1701
		2D6 1719
		2D6 1743

Fig 10B

AMPLICON	LENGTH	SNP ID
		2D6 1757
		2D6 2502
		2D6 2576
		2D6 2593
		2D6 2603
		2D6 2616
		2D6 2642
		2D6 2658
		2D6 3278
		2D6 3280
		2D6 3323
		2D6 3326
		2D6 3343
		2D6 3368
		2D6 3377
		2D6 3465
		2D6 3477
		2D6 3488
		2D6 3562
		2D6 3592
		2D6 3595
		2D6 3597
		2D6 3598
		2D6 4089
		2D6 4099
		2D6 4102
		2D6 4168
		2D6 4194
		2D6 4206
		2D6 4232
		2D6 4469
		2D6 4472
		2D6 4554
		2D6 4557
		2D6 4558
		2D6 4802

AMPLICON	LENGTH	SNP ID
		2D6 4817
		2D6 4896
		2D6 4907
		2D6 5447
		2D6 5472
		2D6 5496
		2D6 5506
		2D6 5661
		2D6 5734
		2D6 5799

Fig 10C

Fig 11 The CodeLink™ SNP Bioarray for human cytochrome P450 genes.

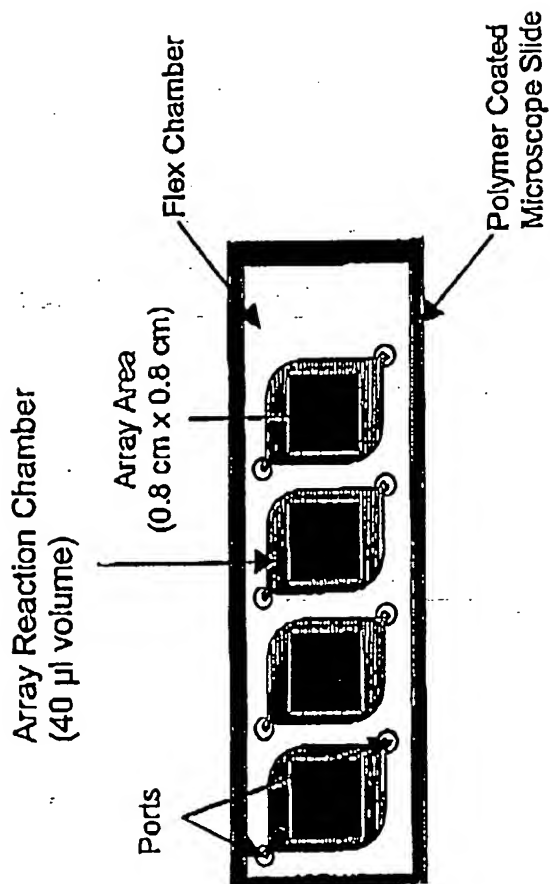
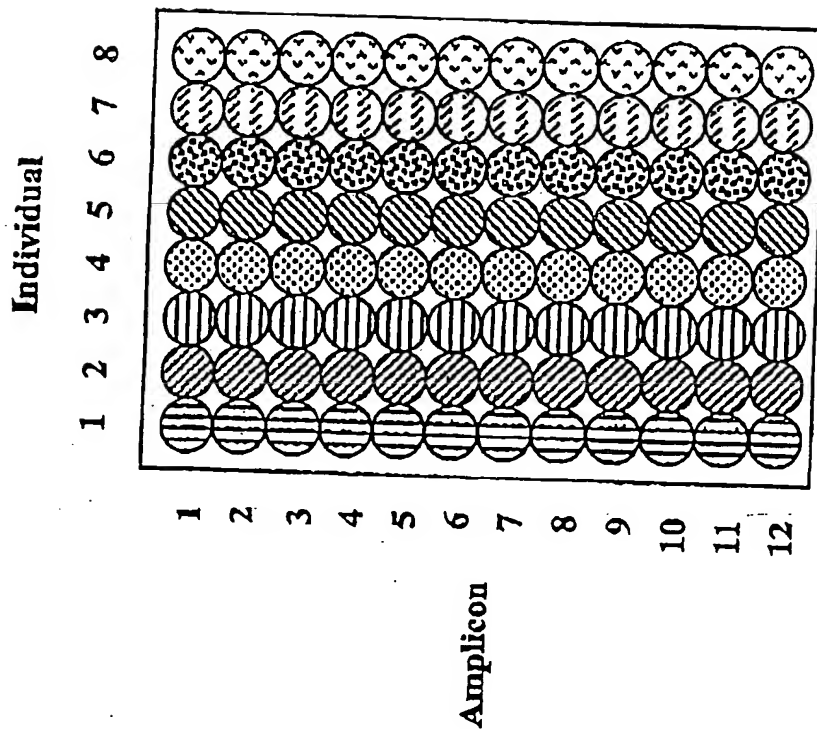


FIG 1 Layout of primer pairs in primer plates.



NOTE: Each sample individual will be aliquoted across 12 separate PCR reactions.

Fig 13. Addition of master mix and samples to microfuge tubes.

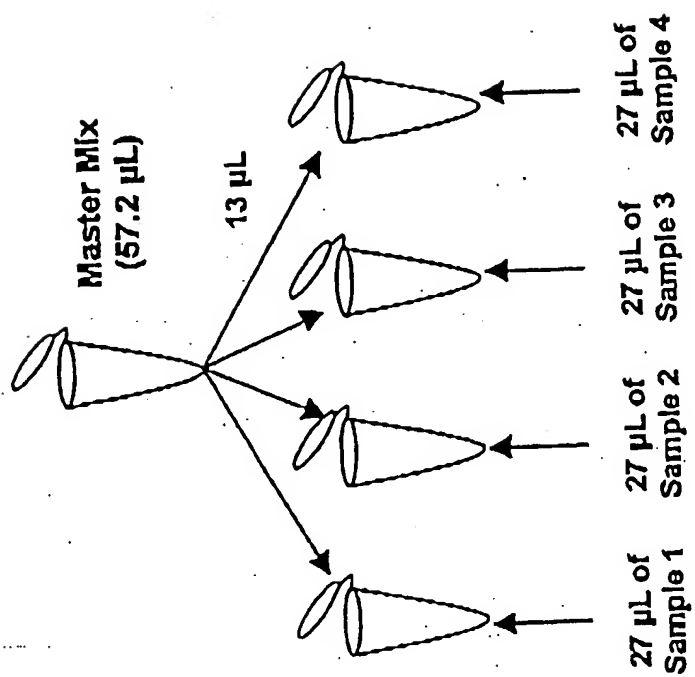


Fig 14 Tip orientation for loading reaction mixtures into chambers.

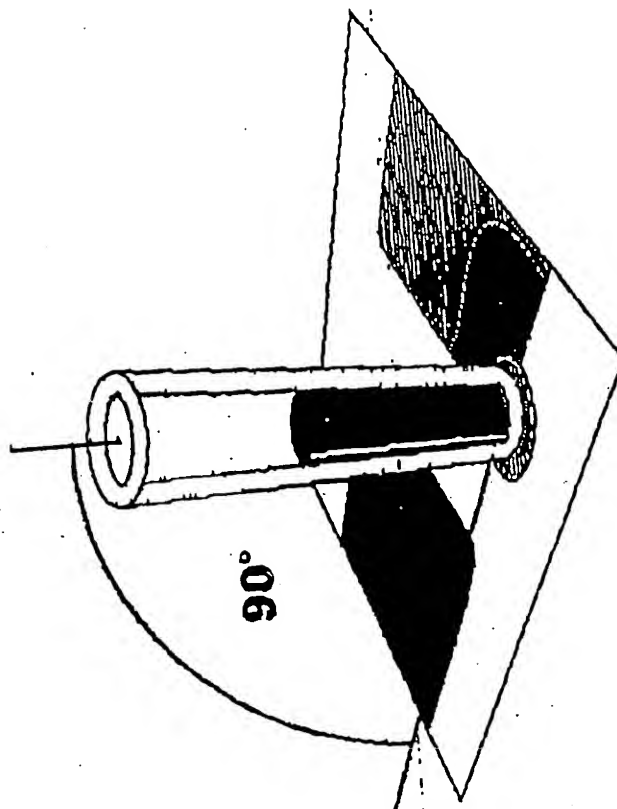


Fig 15 Orientation of sealing strips over chamber ports.

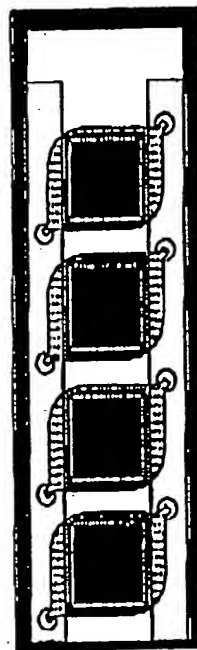
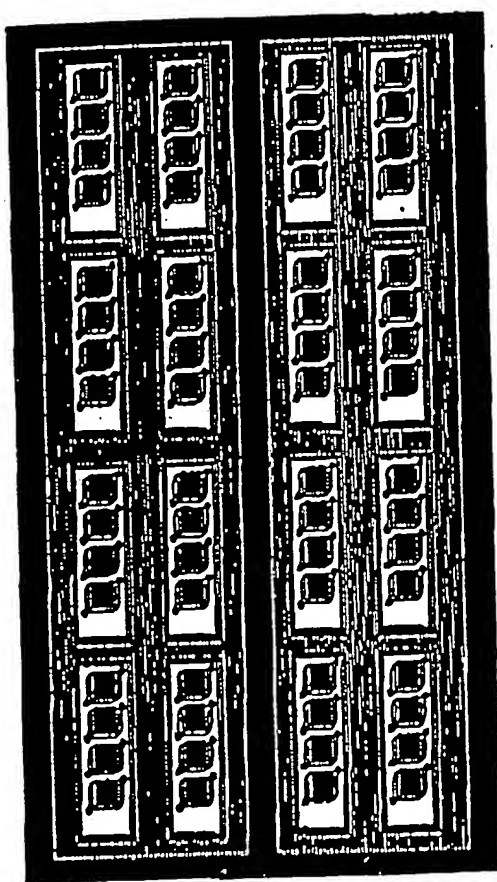


Fig 16 Slide placement on the Hybald Omnislide heat
blocks.



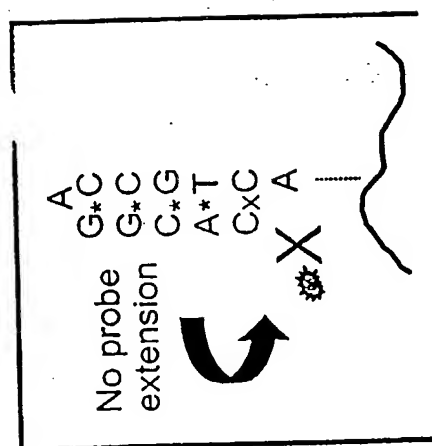


Fig 17

Fig 17 : Prevention of self-extension due to base additions.

Target-Independent signal	Probe	Probe Sequence	Length/#GC	Predicted Stem-Loop	Predicted Base Extended	Observed Base Extended
Strong	504POE321.TA	(5' to 3') TACAGTGGCAGGCA	14/8	<u>TGGCAGGCA</u>	G	G
Strong	60WAF913.114.TA	TCTCTGTCTGTCTCTTGGCA	20/10	<u>TGCTCTTGGCA</u>	G	G
Strong	80POMC07111G.111.CA	AAGTGTCTGATGGAGTAGGAG	21/11	<u>CTGC (8) GGC</u>	C	C
Strong	70POMC07111G.111.CS	GGCAGGCGCAAGGCG	15/12	<u>GGG (8) GGC</u>	G	G
Strong	70LPL2.150.CA	CCGAGATGCTCAGCAGGCTG	21/13	<u>CAGGCTG</u>	G	G
Strong	60WAF288.173.CS	GGCAGGCAATTTATTTC	19/8	<u>GCAG (6) TTTGC</u>	C	C
Strong	APCE182AA	CAGCGCGGCGCT	12/10	<u>GGCGGCGGCT</u>	T	G-C

Table 1 Examples of probe sequences that show a strong target-independent signal in the SBE assay. The predicted stem-loop region is underlined.

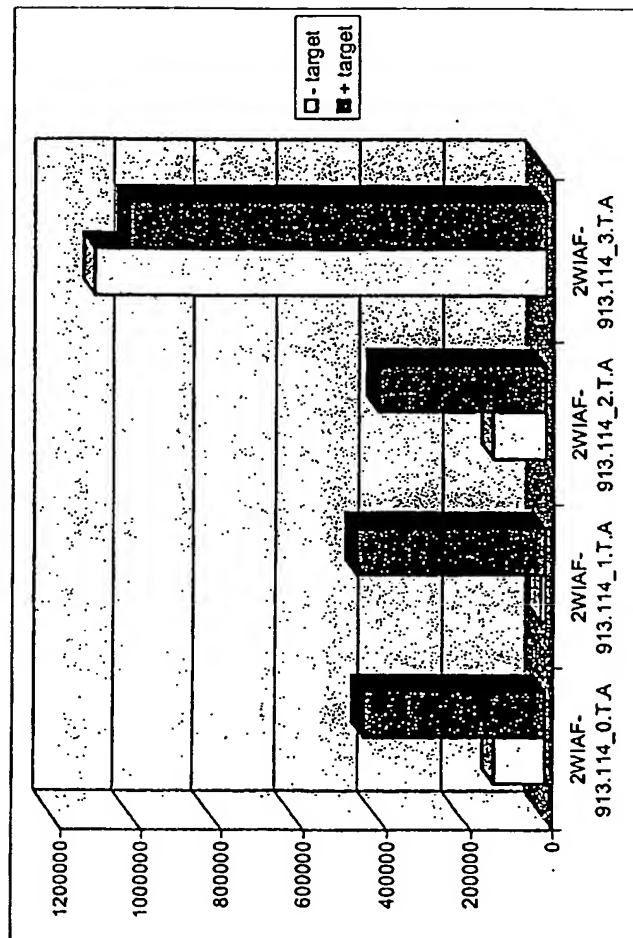
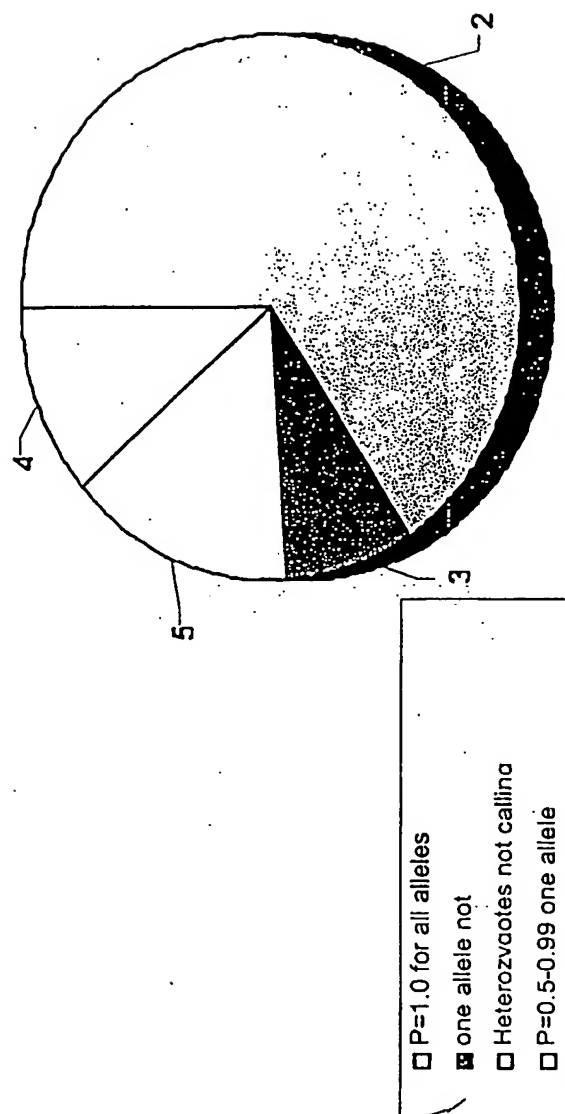


Fig 19

Fig 20



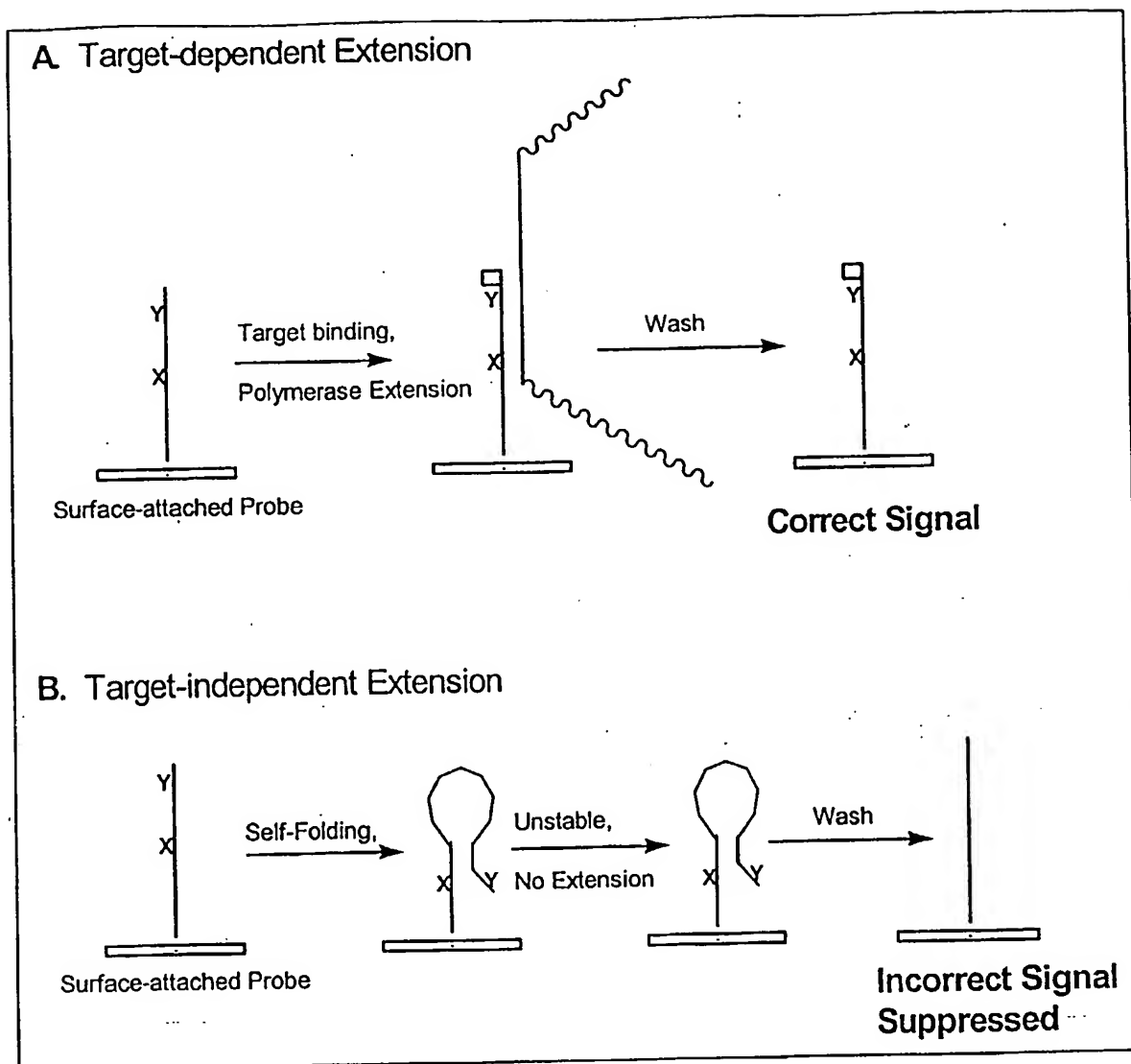
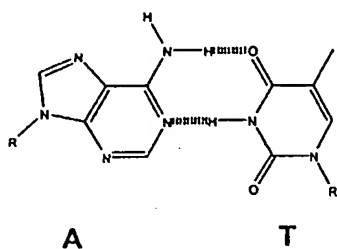
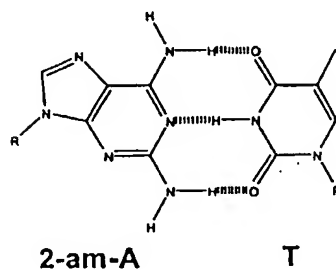


Fig 21

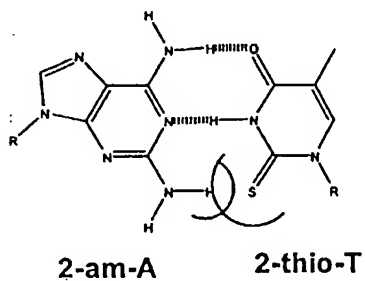
A. Natural A:T base-pair,
pairs equally well with target and itself



C. 2-am-A:T base-pair, (target-probe pair)
forms a very stable base-pair



B. Non-natural 2-am-A:2-thioT base-pair,
does not form a stable base-pair



D. A:2-thio-T base-pair, (target-probe pair)
forms a stable base-pair

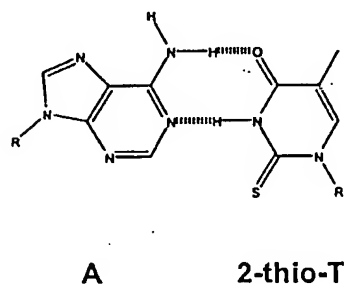
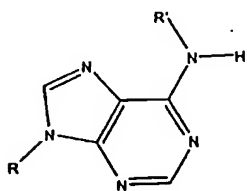


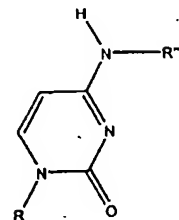
Fig 22

A. Exo-cyclic amine modified A



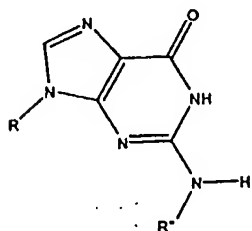
Modified A

A. Exo-cyclic amine modified C



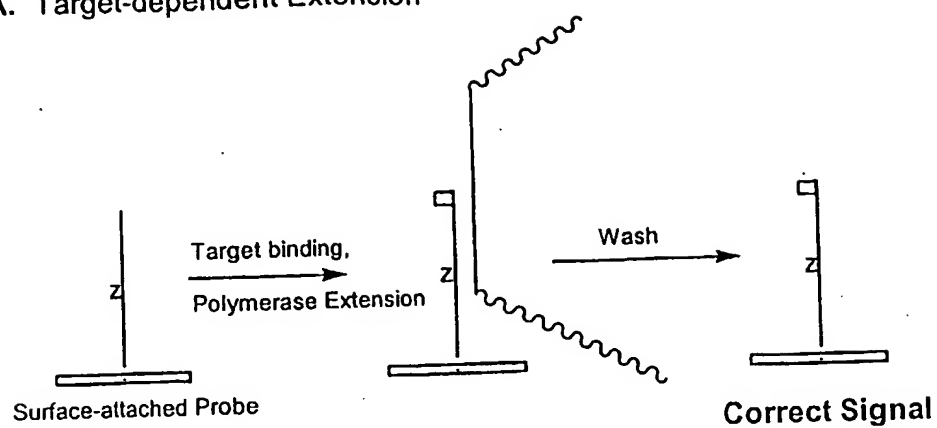
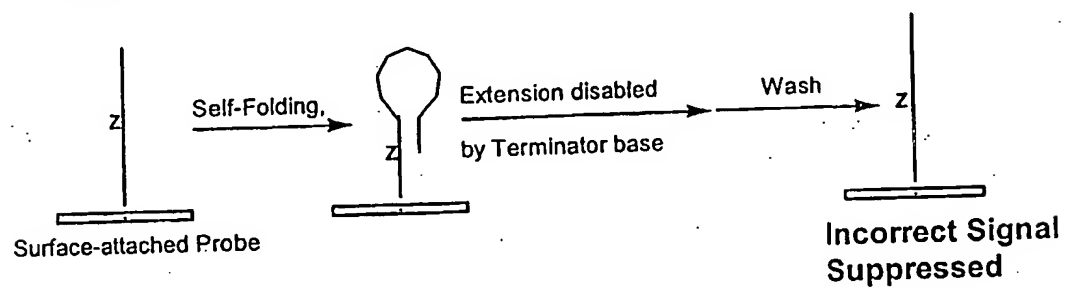
Modified C

B. Exo-cyclic amine modified G

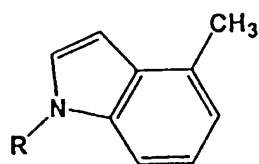


Modified G

Fig 23

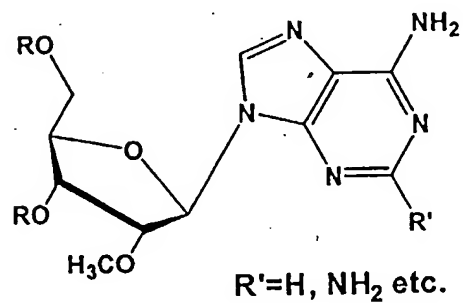
A. Target-dependent Extension**B. Target-independent Extension****Fig 24**

A. "Terminator" base



4-methyl-indole

B. "Terminator" nucleoside



2'-O-methyl-2-amino-A

Fig 25

Fig 26

Table 1 List of modified bases/nucleosides used.

Q = abasic nucleotide; no base-pairing ability, no stacking energy; placed immediately downstream of putative stem-loop; expect A to be incorporated when Q is in template (the "A rule").

I = 4-methylindole; A analog; placed immediately downstream of putative stem-loop; terminates DNA polymerase activity.

K = 5-nitroindole; universal base; placed immediately downstream of putative stem-loop; does not form base pairs but contributes stacking energy.

Z = 2-amino-A; placed within the stem of putative stem-loop; forms 3 hydrogen-bonds with T; no base pairing with 2-thio-dT.

X = 2-thio-dT; placed with stem of putative stem-loop; stable base pairs with A; no base pairing with 2-amino-A.

Fig 26

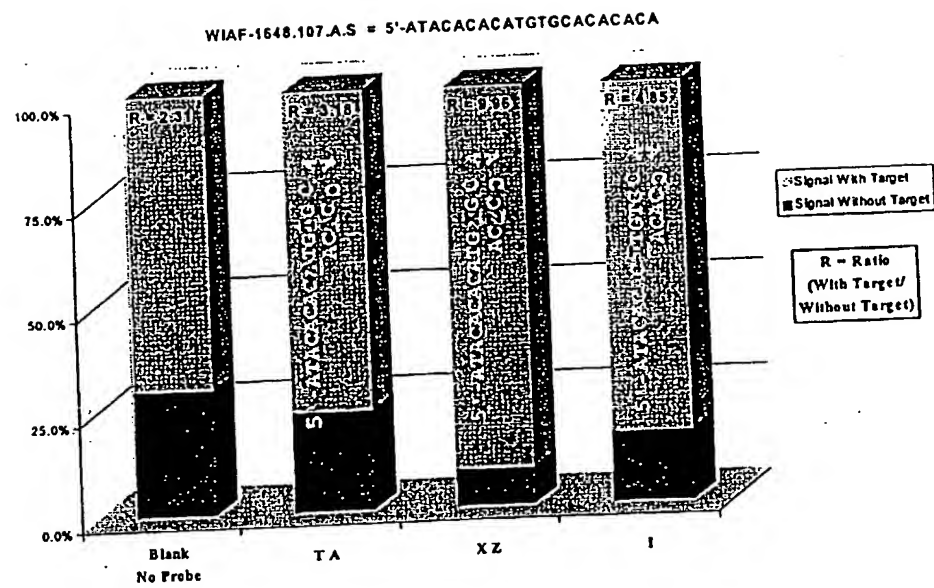


Fig 27

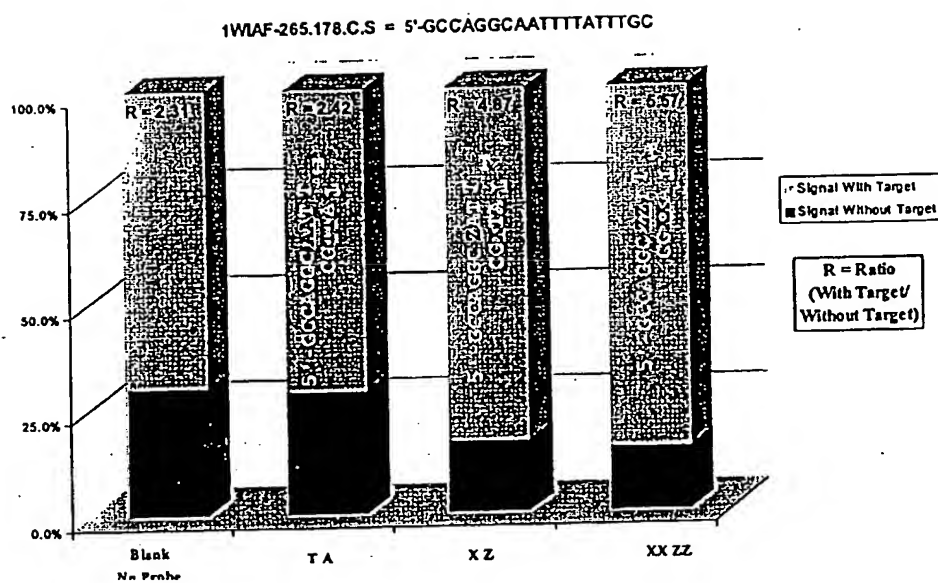


Fig 28

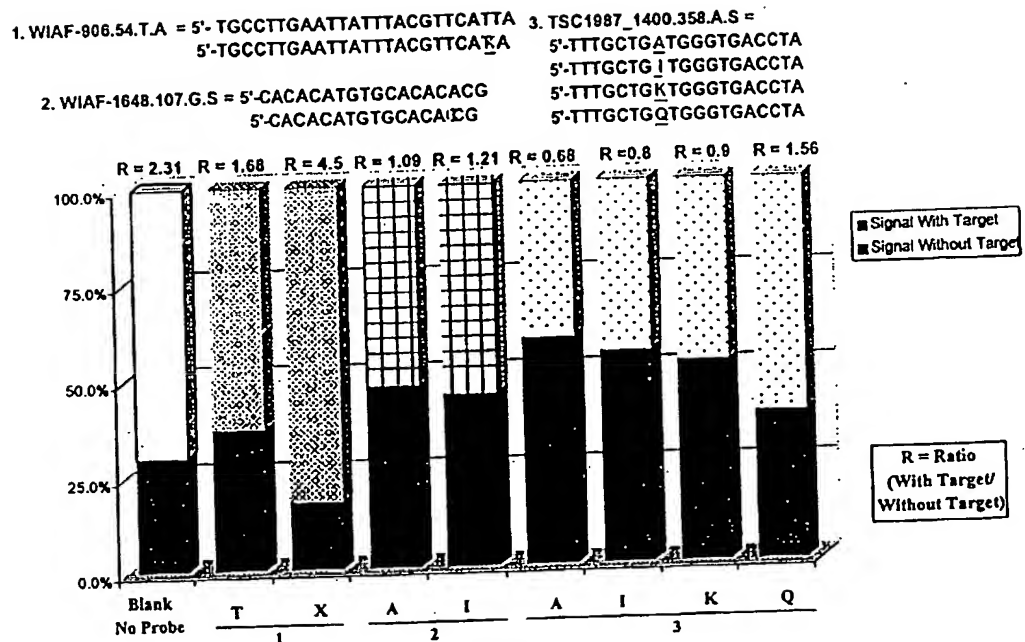
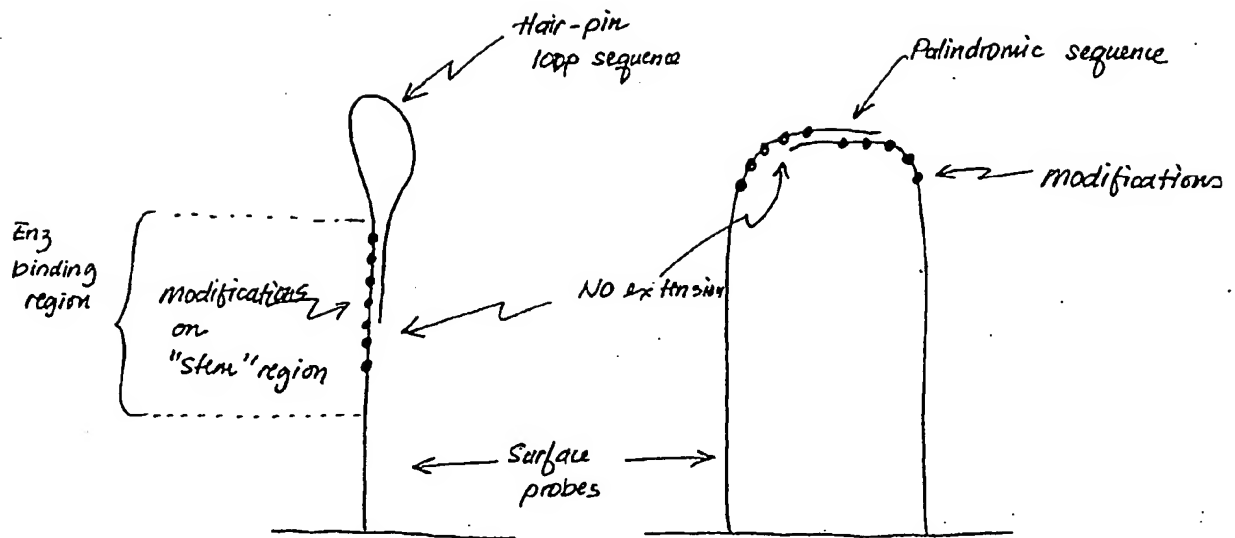
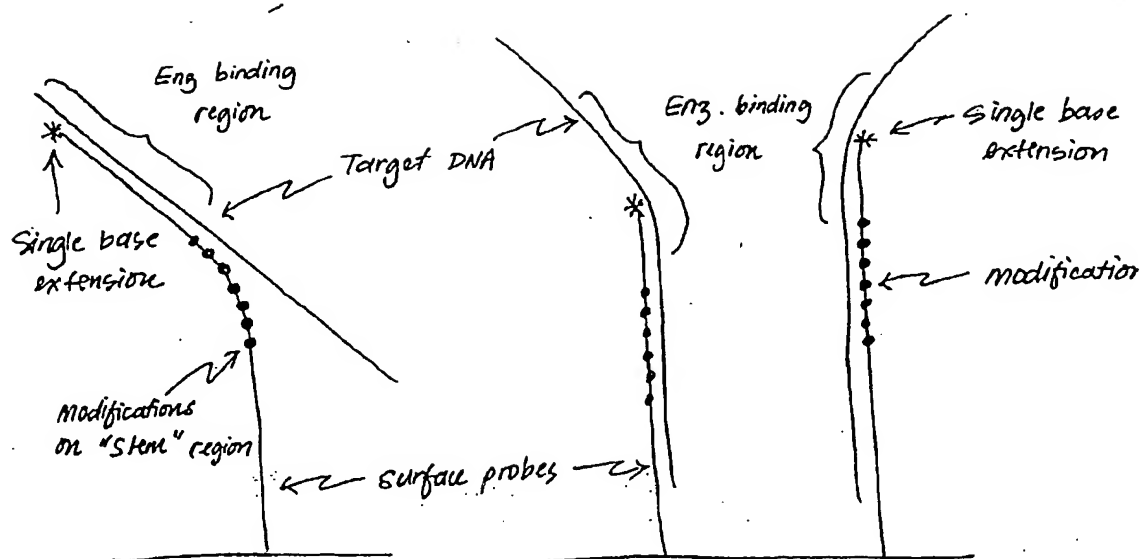


Fig 29

**Fig 30**

Self-Extension inhibited due to modifications on bases in the "stem" region that prevent extension enzyme from binding.

Fig 31



Extension due to probe - target hybrid
is not inhibited.

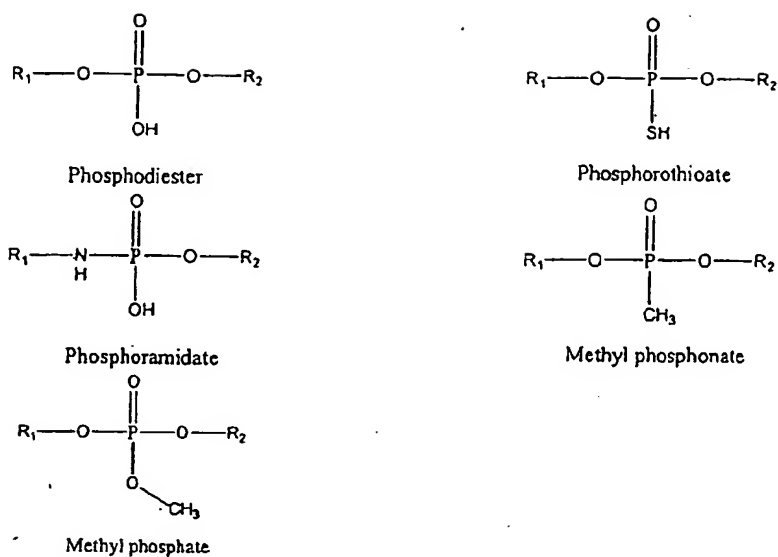


Fig 32A modified nucleotide bases reduces the binding affinity of the SBE enzyme or extension enzyme

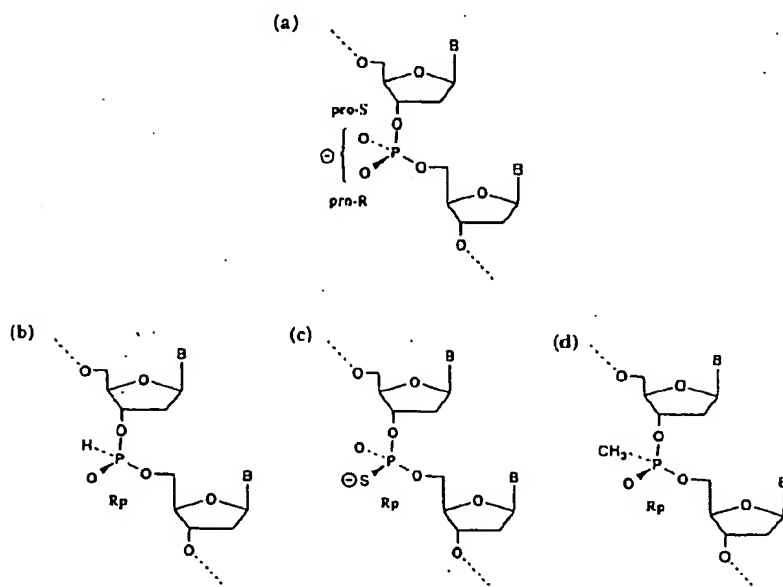


Fig 32B

Chiral phosphodiester analogues

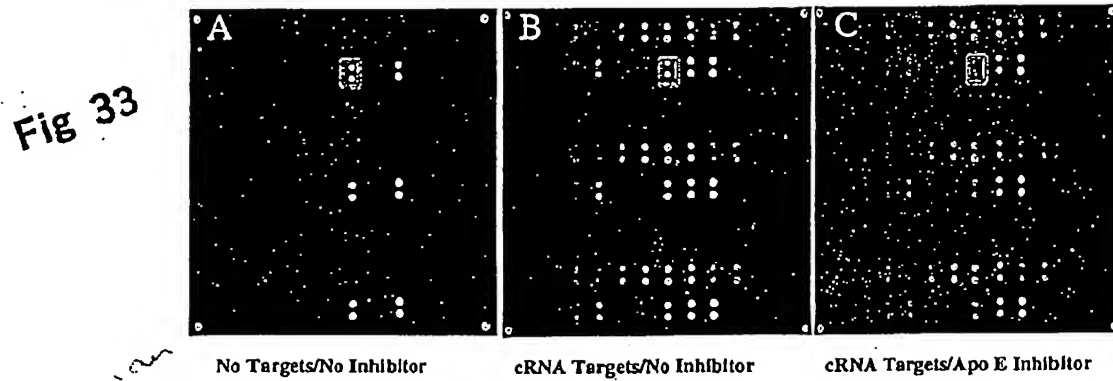


Fig 33 Results from an experiment using oligonucleotide inhibitors to prevent self extension of probes in the SBE array.



Method for uniplexed target prep for SNP genotyping and primer extension without self-extension

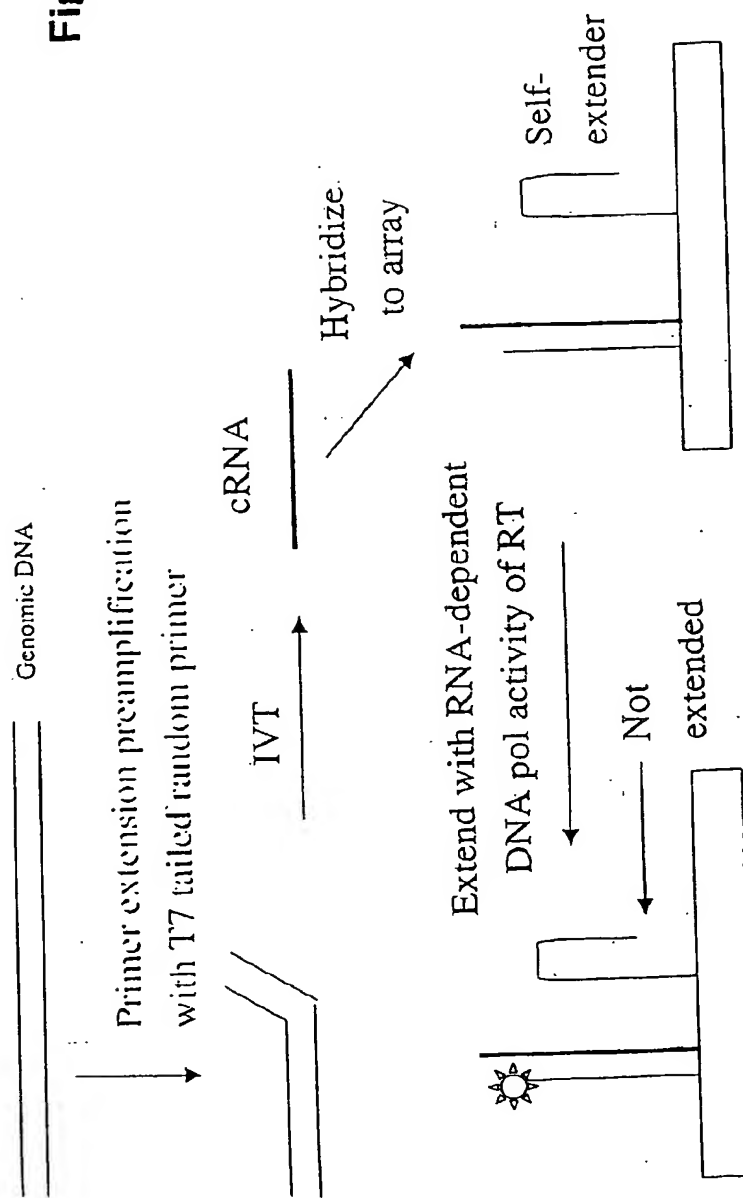


Fig 34

Fig 35

Controls

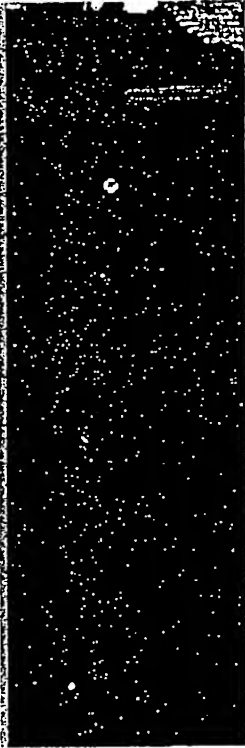
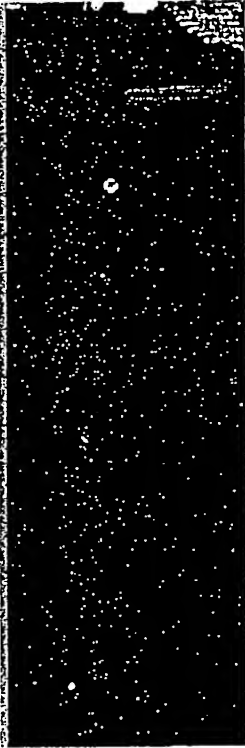
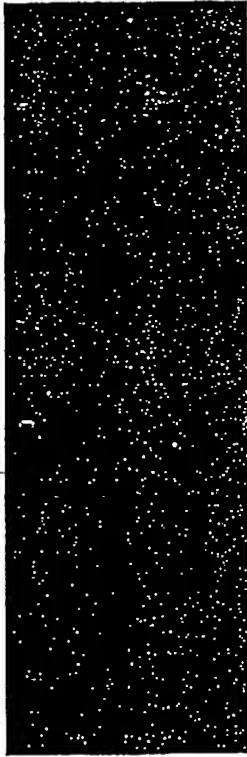
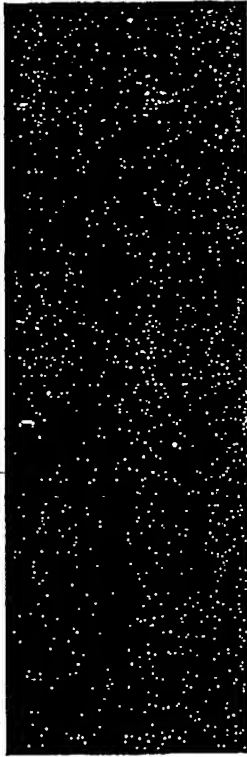
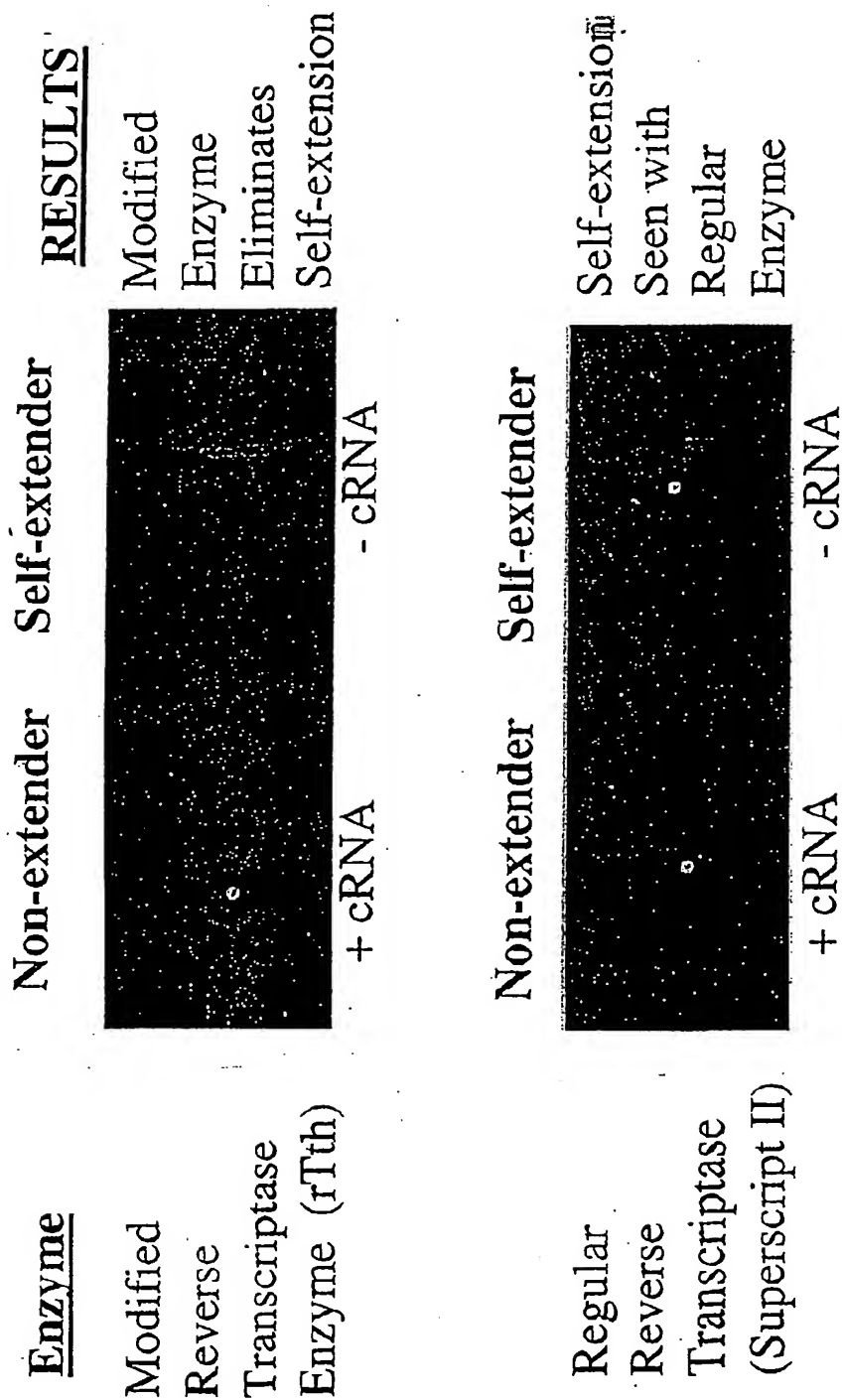
<u>Enzyme</u>	Controls		<u>RESULTS</u>
	Non-extender	Self-extender	
rTth, Modified Reverse transcriptase			The self-extender shows signal only in presence of cRNA
	- cRNA	+ cRNA	
			The signal is human specific since bacterial probes do not show signal with the human cRNA
	Bacterial probe - cRNA	Bacterial probe + cRNA	

Fig 36



Reaction of Amino Oligonucleotides on the SurModics Surface

(an example of acyl substitution reaction on the polymer backbone)

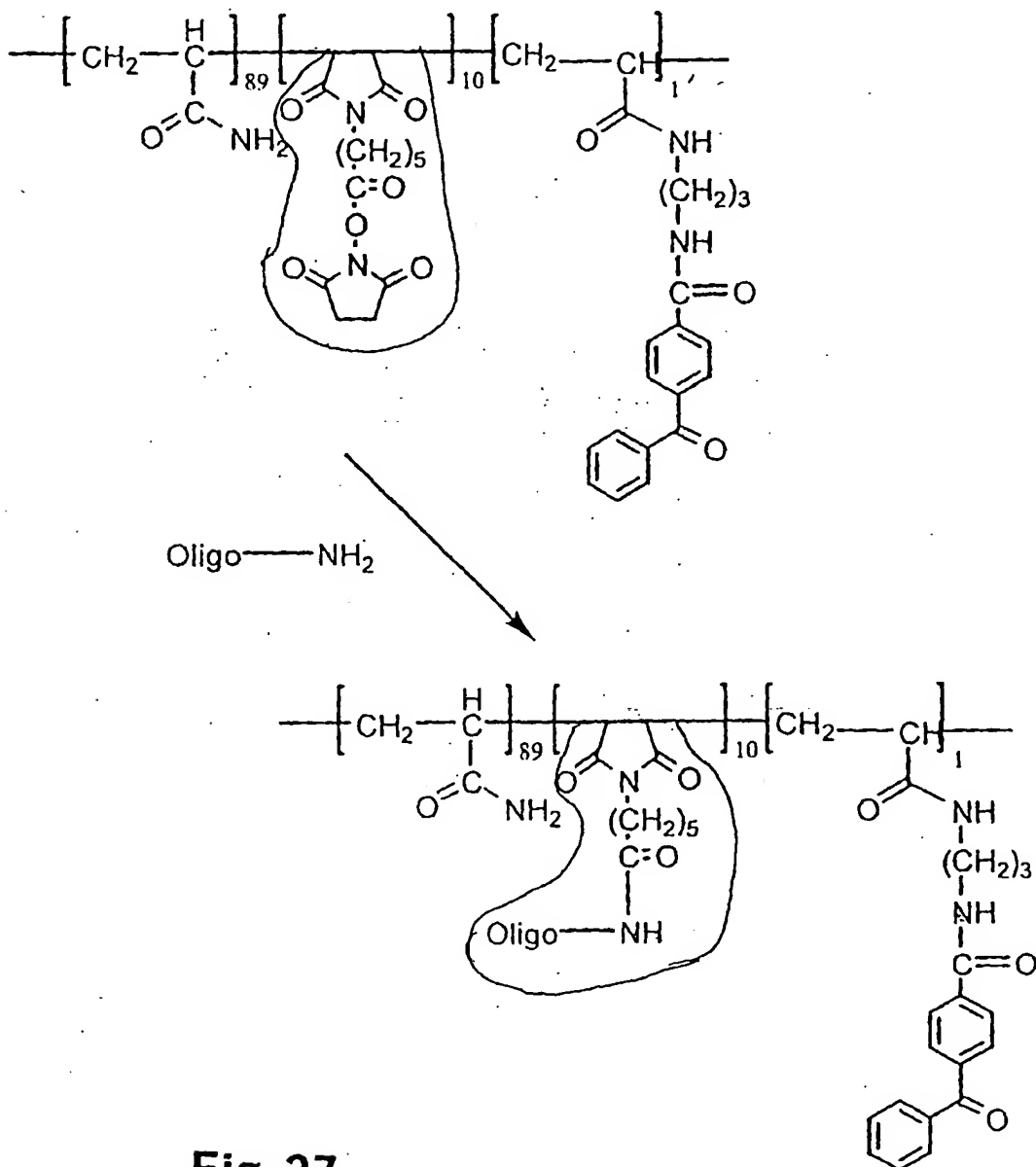
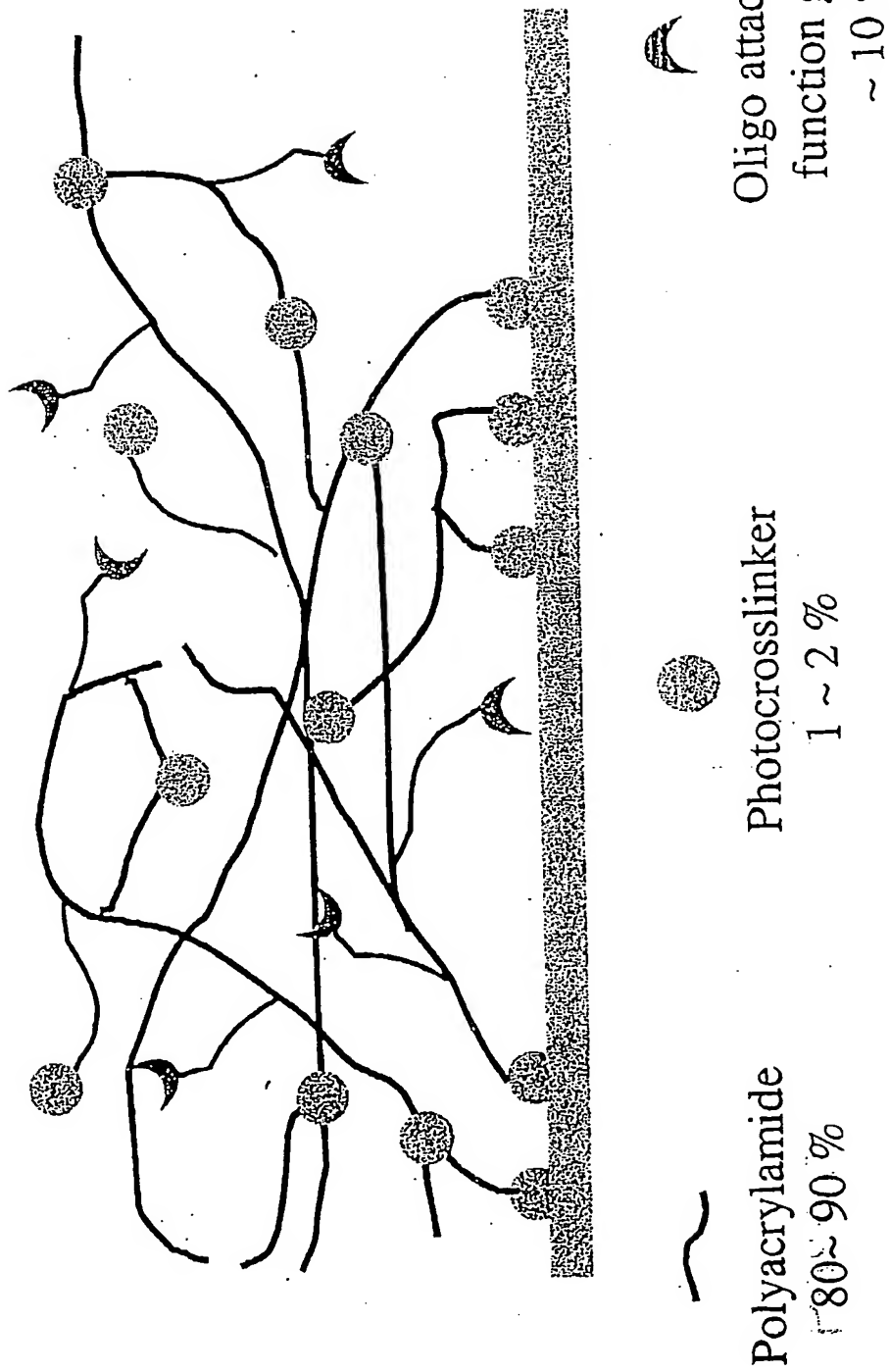


Fig 37

Possible Structure of a SurModics Gel Matrix

**Fig 38**

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